Tape Augmentation Does Not Affect Mid-Term Outcomes of Medial Patellofemoral Ligament Reconstruction in Skeletally Mature Adolescent Patients

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Purpose: To evaluate mid-term outcomes after medial patellofemoral ligament (MPFL) reconstruction with and without tape augmentation in the skeletally mature adolescent population. Methods: All patients under age 18 with recurrent patellar instability treated with surgery at a single institution by a single surgeon from January 2013 through June 2017 were identified by current procedural terminology codes. Inclusion criteria were (1) primary MPFL reconstruction, (2) minimum 3 years’ follow-up, (3) skeletal maturity. Exclusion criteria were (1) bilateral MPFL reconstruction using different techniques on each knee, (2) prior surgery for patellar instability. Chart and imaging review was completed. Patients were contacted to complete a questionnaire, which included the International Knee Documentation Committee (IKDC) form. Results: Fifty-one of 92 eligible patients completed questionnaires. Two patients were excluded. Twenty patients underwent 23 non-augmented MPFL reconstructions; 29 patients underwent 33 augmented MPFL reconstructions. Group demographics were similar. At 4.9 ± 1.2 years follow-up, mean IKDC scores were 77.4 and 79.4 in the nonaugmentation and augmentation groups, respectively. Significantly fewer patients in the augmentation group experienced further injury to their ipsilateral knee compared to the non-augmentation group (6% vs 30%, P = .019). Fewer knees in the augmentation group developed recurrent subjective instability or dislocation after initial surgery requiring surgical correction compared to knees in the nonaugmentation group, although this difference was not significant (6% vs 17%, P = .181). Overall patient-reported outcomes were similar between the 2 groups. Conclusions: There were no significant differences in patient-reported outcomes after MPFL reconstruction with or without tape augmentation. Tape augmentation significantly decreased the risk of subsequent ipsilateral knee injuries, although it did not show a significant difference in recurrent dislocations. Level of Evidence: IV, therapeutic case series.

Acute patellar dislocations account for 2% to 3% of all knee injuries.1 In children younger than age 16, the incidence of patellar dislocation is 43 per 100,000.2–5 Recurrence of patellar dislocation has been reported to occur in 44% to 71% of first time dislocators who do not undergo surgery.6–10 Similarly, an estimated 15% to 44% of patients with first-time patellar dislocations subsequently develop recurrent patellar instability.5,11–16 Although patellar instability may manifest as recurrent patellar dislocations, adolescents and adults may also present without dislocation, as a sense of discomfort, unease, or “something being out of place” in the knee.17

The cause of patellar instability is multifactorial. Several anatomic factors including hypermobility, patella alta, increased tibial tubercle-trochlear groove distance, patellar tilt, rotational deformity, and trochlear dysplasia are prevalent in patients with a history of patellar dislocation or instability.14,18–22 There also appears to be a genetic component, with approximately 28% of patients reporting a family member with
a history of similar symptoms. Patients with recurrent patellar instability have poor outcomes in the absence of surgical intervention. Functional limitations may include pain, mechanical symptoms, or activity restriction, including inability to participate in sport. Long term, patients may worsen chondral damage affecting the patellofemoral joint.

The medial patellofemoral ligament (MPFL) is the primary ligamentous stabilizer that prevents lateral patellar dislocation. MPFL reconstruction for recurrent patellar instability has become a mainstay of operative management for patellar instability. However, with any operative intervention for patellar instability, failure rates have been reported from 5% to 30%, broadly because of technical failure, unaddressed static and dynamic pathoanatomy, or intrinsic risk factors such as collagen disorders or ligamentous laxity. More recently, MPFL reconstruction has been performed using suture augmentation to strengthen the graft and prevent against graft failure. However, the use of synthetic materials has the potential to overconstrain the patellofemoral joint, resulting in prolonged rehabilitation time, anterior knee pain, and degenerative changes of the patellofemoral joint.

The purpose of this study is to evaluate mid-term outcomes after MPFL reconstruction with and without tape augmentation in the skeletally mature adolescent population. We hypothesized that patients in both groups would have improved outcomes, but those who underwent tape augmentation would have fewer ipsilateral re-injuries and surgeries.

Methods

Cohort Selection

A retrospective cohort review was performed with approval from the institutional review board. All patients under age 18 who underwent MPFL reconstruction by the senior author (S.K.A.) from January 2013 through June 2017 were identified by current procedural terminology codes. Inclusion criteria were (1) primary MPFL reconstruction, (2) minimum 3 years’ follow-up, and (3) skeletal maturity. Exclusion criteria were (1) bilateral MPFL reconstruction using different techniques on each knee and (2) prior surgery for patellar instability. All patients underwent a trial of nonoperative management before undergoing surgery. Nonoperative management included activity modification and physical therapy, followed by a gradual progression of activities. Surgery was considered in patients who experienced persistent instability and were unable to return to their desired level of function.

Operative and clinic notes were reviewed to verify index procedure. When available, radiographs were reviewed by 2 fourth-year orthopaedic surgery residents and a medical student (T.E.H., N.J.Q., K.M.T.) for radiographic review of the MPFL reconstruction. When available, radiographs were reviewed by 2 fourth-year orthopaedic surgery residents and a medical student (T.E.H., N.J.Q., K.M.T.) for radiographic review of the MPFL reconstruction.

Surgical Technique and Postoperative Protocol

All patients included in the study underwent MPFL reconstruction with the use of allografts. Patients were placed supine, and general anesthesia was utilized for all procedures. After administration of prophylactic antibiotics, a diagnostic knee arthroscopy was performed to evaluate the chondral surfaces of the patellofemoral joint, remove any loose bodies, and inspect the remainder of the knee for any concomitant pathology that was addressed accordingly. After this, attention was then directed to the open portion of the procedure. Incisions were made over the medial patella and medial epicondyle. Dissection was carried down to the level just above the capsule, where we then tunneled between the two incisions. After exposing the medial patellar surface, a guide pin was placed under fluoroscopic guidance at the junction between the superior third and inferior two thirds of the patella. Once positioning was confirmed, the patellar tunnel was drilled using a 5 mm reamer. The graft, with or without tape augmentation, was trimmed to fit 5 mm tunnels and then fixed into the tunnel using a 4.75 mm PEEK tenodesis interference screw (Arthrex, Naples, FL). When tape augmentation was used, suture tape was whipstitched to the allograft.

Attention was then directed to the femur. A perfect lateral of the knee was obtained using fluoroscopy. A guide pin was then placed slightly proximal and posterior to the medial epicondyle, correlating with Schottle’s point on fluoroscopy. After confirming isometry using the sutures attached to the patellar side, a 5 mm reamer was used on the femoral side, and the graft was fixed with the knee flexed. The flexion angle during fixation was adjusted according to the dynamic exam and findings during isometry testing. The angle chosen for fixation correlated to the position where the graft was the longest, in order to avoid over tensioning. Isometry was checked once again to assure that the graft construct was either isometric or loosened in flexion to confirm that the graft was not over tensioned. Knee range of motion was checked to ensure the patient could achieve full flexion and extension. Wounds were thoroughly irrigated and closed in a layered fashion and patients were placed in a knee immobilizer. Patients were able to bear weight as tolerated using crutches with the knee in full extension beginning the day after surgery. Range of motion was also allowed from 0° to 90° while not weightbearing beginning the day after surgery, with gradual progression as tolerated beginning after the first week.

Survey Methodology

Patient contact information including mailing address, phone number, and e-mail were obtained through review of electronic medical records. Patients
were first contacted by mail alerting them of the study. Patients were then contacted at least 5 times by phone and by e-mail between March 11, 2020, and September 30, 2020, to maximize response rates. Those willing to participate were asked to complete a questionnaire using the Research Electronic Data Capture (REDCap, Vanderbilt University, Nashville, TN) online service. Questionnaires were completed over the phone with the researcher inputting responses directly, or patients were sent an email link to the REDCap survey for completion at a later time. Patients who requested to complete the survey by email were sent automated emails weekly for 5 weeks or until questionnaire completion.

The REDCap questionnaire included the International Knee Documentation Committee (IKDC) form, the Marx Activity Scale, and questions pertaining to knee pain and function, satisfaction with surgery, and additional injury or surgery on either knee (Appendix 1). Chart review was performed to collect demographic and surgery information and to confirm patient-reported additional injuries and surgeries on both knees. Patients were directed to respond to the survey using their worse knee as a reference point.

### Statistical Analysis

Survey data was exported to Microsoft Excel. Data was analyzed using Microsoft Excel and SPSS version 26. All variables were assessed for normality using the Shapiro-Wilk test. Unpaired t-tests were used for normally distributed data and Mann-Whitney U tests were used for non-normally distributed data. Categorical variables were assessed using Pearson $\chi^2$ and Fisher’s exact tests. Significance was set at a $P$ value < .05.

### Results

Eighty-two patients underwent 92 MPFL reconstructions between January 2013 and June 2017. Fifty-one patients (61%) who underwent 59 MPFL reconstructions responded to the survey questionnaire. One patient was excluded because of prior rotational osteotomy for patellar instability, and 1 patient was excluded for bilateral surgery using different techniques on each knee. Among the 49 included patients, 20 patients underwent 23 MPFL reconstructions in the non-tape augmentation group, whereas 29 patients underwent 33 MPFL reconstructions in the tape augmentation group.

Demographics of the 2 groups were similar (Table 1). Mean age, sex distribution, and body mass index were similar at 15.3 years (R: 12.4-17.8), 61% female, and 24.8 kg/m² (R: 16.1, 38.4) versus 15.4 years (R: 12.6-18.2), 70% female, and 24.8 kg/m² (R: 18.1, 37.7) in the nonaugmentation and augmentation groups, respectively. Insall-Salvati ratio, Caton-Deschamps index, and trochlear Dejour classification did not differ

### Table 1. Patient Demographics

|                                | Nonaugmentation (n = 23) | Augmentation (n = 33) | $P$ value |
|--------------------------------|--------------------------|-----------------------|-----------|
| Age at time of surgery, y      | 15.3 (1.6)               | 15.4 (1.6)            | .782      |
| Mean (SD)                      | 12.4, 17.8               | 12.6, 18.2            |           |
| Sex                            |                          |                       |           |
| Female                         | 14 (61%)                 | 23 (70%)              | .492      |
| Male                           | 9 (39%)                  | 10 (30%)              |           |
| Operative knee                 |                          |                       | .299      |
| Left                           | 13 (57%)                 | 14 (42%)              |           |
| Right                          | 10 (43%)                 | 19 (58%)              |           |
| BMI, kg/m²                     |                          |                       | .298      |
| Mean SD                        | 23.7 (5.3)               | 24.8 (4.8)            |           |
| Range                          | 16.1, 38.4               | 18.1, 37.7            |           |
| Insall Salvati ratio,* mean (SD)| 1.3 (0.3)               | 1.4 (0.3)             | .735      |
| Caton Deschamps index,* mean (SD)| 1.3 (0.2)              | 1.3 (0.2)             | .981      |
| Trochlear dysplasia, n (%)     |                          |                       | .699      |
| No                             | 5 (22%)                  | 14 (42%)              |           |
| Yes                            | 7 (30%)                  | 15 (45%)              |           |
| Trochlear Dejour classification* |                        |                       | .267      |
| A                              | 6 (26%)                  | 14 (42%)              |           |
| B                              | 0 (0%)                   | 1 (3%)                |           |
| C                              | 1 (4%)                   | 0 (0%)                |           |
| D                              | 0 (0%)                   | 0 (0%)                |           |
| Preoperative IKDC              |                          |                       | .718      |
| Mean (SD)                      | 46.8 (16.8)              | 40.3 (18.1)           |           |
| Range                          | 33.3, 70.1               | 6.9, 69.0             |           |

* Lateral films were unavailable in 3 knees (13%) in the nonaugmentation group and 1 knee (3%) in the augmentation group. Sunrise view radiographs were unavailable in 11 knees (48%) in the nonaugmentation group and 4 knees (12%) in the augmentation group.

$^{1}$ Preoperative IKDC scores were available in 4 knees (17%) in the nonaugmentation group and 23 knees (70%) in the augmentation group.
significantly between the 2 groups. Lateral films were unavailable in 3 knees (13%) in the nonaugmentation group and 1 knee (3%) in the augmentation group. Sunrise view radiographs were unavailable in 11 knees (48%) in the nonaugmentation group and 4 knees (12%) in the augmentation group.

Surgical data are described in Table 2. The augmentation group underwent more procedures than the nonaugmentation group at the time of surgery (58% vs 22%, \(P = .008\)), with 15 of 19 patients (45%) in the augmentation group undergoing loose body removal.

Patient reported outcomes are detailed in Table 3. The augmentation group had shorter follow-up at a mean of 4.1 years (R: 3.0-5.2) versus 6.0 years (R: 4.2-7.0) in the nonaugmentation group (\(P < .001\)). Outcomes did not differ significantly between the two groups. Mean IKDC scores were 79.4 and 77.4 in the augmentation and non-augmentation groups, which exceeded the patient acceptable symptom state (PASS) previously defined as 75.9.\(^{34,35}\) Mean Marx activity scale scores were 8.6 and 9.2, respectively. The augmentation group reported their operative knee as 82.4% (R: 40-100) of “normal,” whereas the nonaugmentation group reported their operative knee as 80.9% (R: 17-100) of “normal.”

Pain ratings did not differ significantly between the two groups. Regarding pain on a visual analog scale (VAS), the percentages of patients reporting pain less than a 3 at rest, with daily activity, and with sport were 86% vs 85% (\(p = 1.000\)), 72% vs. 70% (\(p = 0.915\)), and 45% vs 55% (\(p = 0.484\)) in the augmentation and non-augmentation groups, respectively.

Satisfaction with surgery as rated on a Likert scale did not differ between the 2 groups (\(P = .305\)). Overall patient satisfaction was high in both groups. Ten patients (50%) reported being “very satisfied” in the nonaugmentation group, compared with 19 patients (66%) in the augmentation group. When asked whether they would pursue MPFL surgery again, 18 patients (90%) in the nonaugmentation group responded “definitely yes,” compared with 23 patients (79%) in the augmentation group.

Patients who underwent MPFL reconstruction participate in sport at similar levels as prior to surgery, regardless of technique used (31% augmentation vs 20% nonaugmentation, \(P = .516\)). Patients in the augmentation group avoid sport more than their counterparts who did not undergo augmentation (41% vs 30%), although this difference was not significant (\(P = .417\)). Of patients who refrained from sport, 75% in the augmentation group versus 83% in the non-augmentation group reported doing so due to their knee (\(P = .646\)).

Ten patients (50%) in the augmentation group reported stiffness or loss of motion in their knee, versus 12 patients (41%) in the nonaugmentation group (\(P = .551\)). Of patients who reported stiffness or loss of motion, 75% of those in the augmentation group and 40% of those in the non-augmentation group reported limitation of activity due to these symptoms (\(P = .192\)). Eight patients (28%) in the augmentation group reported instability occurring weekly or more, compared with 6 patients (26%) in the nonaugmentation group (\(P = .495\)).

Patients in the augmentation group sustained fewer ipsilateral knee injuries than the non-augmentation group (6% vs 30%, \(P = .019\)). Time between initial surgery and subsequent ipsilateral injury was 2.5 ± 2.1 years in the nonaugmentation group, whereas only 1 patient in the augmentation group sustained an ipsilateral injury 2.0 years after surgery. Overall, 2 knees (6%) in the augmentation group developed recurrent subjective instability or dislocation after initial surgery requiring surgical correction, compared to 4 knees (17%) knees in the nonaugmentation group (\(P = .181\)).

In the augmentation group, these injuries included

| Procedure performed                  | Nonaugmentation (n = 23) | Augmentation (n = 33) | \(P\) Value |
|--------------------------------------|--------------------------|----------------------|-------------|
| Isolated MPFL reconstruction         | 18 (78%)                 | 14 (32%)             | .008*       |
| Chondral debridement                 | 2 (9%)                   | 3 (9%)               |             |
| Partial medial meniscectomy          | 0 (0%)                   | 1 (3%)               |             |
| Partial lateral meniscectomy         | 0 (0%)                   | 1 (3%)               |             |
| Lateral meniscus repair              | 0 (0%)                   | 1 (3%)               |             |
| ACL reconstruction                   | 0 (0%)                   | 1 (3%)               |             |
| Loose body removal                   | 3 (12%)                  | 15 (45%)             | <.001*      |
| Graft                                |                          |                      |             |
| Gracilis allograft                   | 2 (9%)                   | 21 (63%)             |             |
| Peroneus longus allograft            | 1 (4%)                   | 0 (0%)               |             |
| Semitendinosis allograft             | 19 (82%)                 | 12 (36%)             |             |
| Tibialis anterior allograft          | 1 (3%)                   | 0 (0%)               |             |

*Significant at the .05 level.
nonspecific knee injury (n = 1) and instability event (n = 1). For the nonaugmentation group, injuries included ACL rupture (n = 1), patellar dislocation (n = 1), instability event (n = 4), and nonspecific knee injury (n = 1). Five knees (22%) within the nonaugmentation group underwent further surgery on the ipsilateral knee, compared with 2 knees (6%) in the augmentation group (P = 0.081). Time between initial and subsequent ipsilateral surgery was 2.2 ± 1.5 years in the nonaugmentation group and 2.3 ± 1.6 years in the augmentation group (P = 0.938). In the augmentation group, subsequent surgeries included revision MPFL reconstruction and tibial tubercle osteotomy (n = 1), and revision MPFL reconstruction with distal femoral lateral opening wedge osteotomy, which was followed by a second revision MPFL reconstruction and hardware removal (n = 1).

### Discussion

In this series, adolescent patients undergoing MPFL reconstruction with and without tape augmentation for patellar instability at a mean of 4.9 ± 1.2 years follow-up had similar patient-reported outcomes. However, fewer patients in the tape augmentation group experienced further injury of their ipsilateral knee. Patients in both groups reported similar levels of recurrent patellar instability, including dislocation.

Patient-reported outcomes, including IKDC score, reported knee percent of normal, and VAS pain scores, did not differ significantly between the augmentation and non-augmentation groups. The results suggest that MPFL tape augmentation may offer additional support for patellar stability in adolescent patients, potentially reducing the risk of further injury and the need for subsequent surgery.
and non-augmentation groups. Xie et al similarly evaluated the clinical outcome of MPFL reconstruction for patellar instability using semitendinosis tendons with and without polyester suture augmentation, but found significant differences in IKDC score between the augmentation and non-augmentation groups, with the augmentation group overall demonstrating better outcomes.

Harris et al. published minimal clinically important difference and PASS scores for commonly used patient report outcome scores and showed that for IKDC, minimal clinically important difference was 16.7, whereas PASS was 75.9. Patients in both groups surpassed both of these thresholds because the change from preoperative to postoperative IKDC scores were 39.1 and 30.6 for the augmentation and non-augmentation groups, respectively; the average postoperative scores for each group was 79.4 and 77.4, respectively.

In a systematic review of pediatric and adult patients undergoing MPFL reconstruction, Manjunath et al. found that the overall return to play rate was 85.1%, with 68.3% returning to the same level of play. Sport participation was lower overall in the current study population. Patients who underwent MPFL reconstruction participate in sport at similar levels, regardless of technique used (20% nonaugmentation vs 31% augmentation). Harris et al. published minimal clinically important difference and PASS scores for commonly used patient report outcome scores and showed that for IKDC, minimal clinically important difference was 16.7, whereas PASS was 75.9. Patients in both groups surpassed both of these thresholds because the change from preoperative to postoperative IKDC scores were 39.1 and 30.6 for the augmentation and non-augmentation groups, respectively; the average postoperative scores for each group was 79.4 and 77.4, respectively.

Manjunath et al. found that the rate of recurrent instability events following MPFL reconstruction in a combined pediatric and adult population was 5.4%. Lind et al. found that 20% of pediatric patients experienced re-dislocation, compared with 5% in the adult population. The current study found that 2 knees (6%) in the augmentation group developed recurrent subjective instability or dislocation after initial surgery requiring surgical correction, compared to 4 knees (17%) in the nonaugmentation group (P = .181), with an overall rate of 11% (6/56). Pediatric patients appear more prone to recurrent instability and dislocation after MPFL reconstruction than their adult counterparts. The data from the current study show that tape augmentation might play a role in improving outcomes in skeletally mature adolescent patients with regard to recurrent instability rates after MPFL reconstruction.

In their comparison between patients who underwent MPFL reconstruction with and without polyester suture augmentation, Xie et al. found that no patients in the augmentation group experienced redislocation, compared with 2 patients (4.7%) in the nonaugmentation group, and that failure occurred in 1 patient (2.4%) in the augmentation group versus 10 patients (23.3%) in the nonaugmentation group. The findings of the current study align with those of Xie et al. Two of 34 knees (5.8%) underwent revision MPFL reconstruction in the augmentation group, compared with 4 of 24 knees (16.7%) in the non-augmentation group. However, the present study differs from that of Xie et al. in that there was a statistically significant increase in the number of injuries sustained to the ipsilateral knee between the nonaugmentation group (29.2%) and the augmentation group (6.3%). Notably, procedures in the present study did not use a standard graft type; gracilis, semitendinosus, tibialis anterior, and peroneus longus allograft were used. However, Stupay et al. demonstrated in their systematic review of 34 articles that graft choice, excluding tape augmentation, did not impact outcomes or complications of MPFL reconstruction, supporting that the observed difference in the present study is likely attributable to tape augmentation.

Graft deterioration may contribute to subsequent ipsilateral injuries in the nonaugmented group. Zhao et al. demonstrated that correction of static patellar position deteriorated over a 5-year follow-up after MPFL reconstruction without polyester suture augmentation. Xie et al. hypothesized that shortly after harvest, the tendon loses blood supply and suffers necrosis, such that the initial tendon strength observed deteriorates over time. Further cadaveric biomechanical study may better elucidate the relationship between graft age and failure load with and without tape augmentation.

**Limitations**

There are several limitations in this study. First, the response rate was 61%. This loss to follow-up leads to attrition bias, which may affect the validity of the conclusions. A larger sample size may have demonstrated more significant differences between the 2 groups. Multiple attempts were made to contact study patients over a several-month period. Second, procedures in the present study did not use a standard graft type; gracilis, semitendinosis, tibialis anterior, and peroneus longus allograft were used. Third, follow-up time differed significantly between the 2 groups. Patients in the augmentation group had shorter follow-up because of tape augmentation not having been incorporated into the primary surgeon’s (S.K.A.) practice until more recently. Fourth, lack of availability of preoperative radiographs limits potential elucidation of morphologic differences that may affect patellar stability, including trochlear dysplasia, between the two groups. Fifth, this study is inherently limited through its design as a retrospective study. Finally, all patients underwent
surgery at a single institution by a single surgeon (S.K.A.), which may limit generalizability of these findings, although use of a single surgeon minimizes discrepancies in surgical technique.

**Conclusion**

There were no significant differences in patient-reported outcomes after MPFL reconstruction with or without tape augmentation. Tape augmentation significantly decreased the risk of subsequent ipsilateral knee injuries, although did not show a significant difference in recurrent dislocations.

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### Appendix 1. Patient Survey

#### Demographics

| Question                                                                 | Options          |
|--------------------------------------------------------------------------|------------------|
| What is your name?                                                       |                  |
| Are you filling this questionnaire out for yourself or your child?       | My self, My child|
| Was surgery performed on the affected knee?                              | Yes, No          |
| On which knee did Dr. XXX or Dr. XXX perform surgery?*                   | Left, Right, Both|

#### 2000 IKDC Subjective Knee Evaluation Form

| Question                                                                 | Options                                                                 |
|--------------------------------------------------------------------------|-------------------------------------------------------------------------|
| What is the highest level of activity that you can perform without significant knee pain? | Very strenuous activities like jumping or pivoting as in basketball or soccer |
|                                                                          | Strenuous activities like heavy physical work, skiing or tennis          |
|                                                                          | Moderate activities like moderate physical work, running or jogging      |
|                                                                          | Light activities like walking, housework or yardwork                    |
|                                                                          | Unable to perform any of the above activities due to knee pain           |
| During the past 4 weeks, or since your injury, how often have you had pain? (0 = Never and 10 = Constant) | 0, 1, 2, 3, 4, 5, 6, 7, 8, 9, 10 |
| If you have pain, how severe is it? (0 = No pain and 10 = worst pain imaginable) | 0, 1, 2, 3, 4, 5, 6, 7, 8, 9, 10 |
| During the past 4 weeks, or since your injury, how stiff or swollen was your knee? | Not at all, Mildly, Moderately, Very, Extremely |
| What is the highest level of activity you can perform without significant swelling in your knee? | Very strenuous activities like jumping or pivoting as in basketball or soccer |
|                                                                          | Strenuous activities like heavy physical work, skiing or tennis          |
|                                                                          | Moderate activities like moderate physical work, running or jogging      |
|                                                                          | Light activities like walking, housework or yardwork                    |
|                                                                          | Unable to perform any of the above activities due to knee pain           |
| During the past 4 weeks, or since your injury, did your knee lock or catch? | Yes, No |

(continued)
What is the highest level of activity you can perform without significant giving way in your knee?

- Very strenuous activities like jumping or pivoting as in basketball or soccer
- Strenuous activities like heavy physical work, skiing or tennis
- Moderate activities like moderate physical work, running or jogging
- Light activities like walking, housework or yardwork
- Unable to perform any of the above activities due to knee pain

What is the highest level of activity you can participate in on a regular basis?

- Very strenuous activities like jumping or pivoting as in basketball or soccer
- Strenuous activities like heavy physical work, skiing or tennis
- Moderate activities like moderate physical work, running or jogging
- Light activities like walking, housework or yardwork
- Unable to perform any of the above activities due to knee pain

How does your knee affect your ability to:

| Activity                          | Not difficult at all | Minimally difficult | Moderately difficult | Extremely difficult | Unable to |
|-----------------------------------|----------------------|---------------------|----------------------|---------------------|-----------|
| a. Go up stairs                   | ○                    | ○                   | ○                    | ○                   | ○         |
| b. Go down stairs                 | ○                    | ○                   | ○                    | ○                   | ○         |
| c. Kneel on the front of your knee| ○                    | ○                   | ○                    | ○                   | ○         |
| d. Squat                          | ○                    | ○                   | ○                    | ○                   | ○         |
| e. Sit with your knee bent        | ○                    | ○                   | ○                    | ○                   | ○         |
| f. Rise from a chair              | ○                    | ○                   | ○                    | ○                   | ○         |
| g. Run straight ahead             | ○                    | ○                   | ○                    | ○                   | ○         |
| h. Jump and land on your involved leg | ○                  | ○                   | ○                    | ○                   | ○         |
| i. Stop and start quickly         | ○                    | ○                   | ○                    | ○                   | ○         |

Function: How would you rate the function of your knee on a scale of 0 to 10 with 10 being normal, excellent function and 0 being the inability to perform any of your usual daily activities which may include sports?

Function prior to your knee injury:
(0 = Cannot perform daily activities and 10 = No limitation in daily activities)

| Rating | 0 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 | 9 | 10 |
|--------|---|---|---|---|---|---|---|---|---|---|----|

(continued)
Current function of your knee: (0 = Cannot perform daily activities and 10 = No limitation in daily activities)

| 0 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 | 9 | 10 |

**Marx Activity Scale**

Please indicate how often you performed each activity in your healthiest and most active state, in the past year.

- **Running**: running while playing a sport or jogging
  - Less than one time in a month
  - One time in a month
  - One time in a week
  - 2 or 3 times in a week
  - 4 or more times in a week

- **Cutting**: changes directions while running
  - Less than one time in a month
  - One time in a month
  - One time in a week
  - 2 or 3 times in a week
  - 4 or more times in a week

- **Decelerating**: coming to a quick stop while running
  - Less than one time in a month
  - One time in a month
  - One time in a week
  - 2 or 3 times in a week
  - 4 or more times in a week

- **Pivoting**: turning your body with your foot planted while playing a sport; For example: skiing, skating, kicking, throwing, hitting a ball (golf, tennis, squash), etc.
  - Less than one time in a month
  - One time in a month
  - One time in a week
  - 2 or 3 times in a week
  - 4 or more times in a week

**Survey**

- **How would you rate your affected knee today as a percentage of normal (0-100% scale with 100% being “normal”)?**
  - 0%
  - 50%
  - 100%

- **Please indicate how often you experienced knee instability events (i.e., the feeling of your knee giving way), in the past year?**
  - Less than one time in a month
  - One time in a month
  - One time in a week
  - 2 or 3 times in a week
  - 4 or more times in a week

- **How satisfied are you with the results of your surgery?**
  - Very Satisfied
  - Satisfied
  - Neutral
  - Unsatisfied
  - Very Unsatisfied

- **Looking back, if you “had to do it all over again”, would you have the**
  - Definitely, yes
  - Probably, yes
| Question                                                                 | Options |
|-------------------------------------------------------------------------|---------|
| Have you had any further surgeries on your knee since your initial knee surgery with Dr. XXX or Dr. XXX?*† | Yes, No |
| Please explain what further surgeries you’ve had since your initial knee surgery with Dr. XXX or Dr. XXX.* Include approximate date of surgery, if known. | Yes, No |
| Since your knee surgery, have you experienced any other injuries to your surgical knee?† | Yes, No |
| Since your knee surgeries, have you experienced any injuries to your other knee?† | Yes, No |
| Do you currently play any sports? | Yes, No |
| What sports do you currently play and at what level (competitive, recreational, etc)? | (Example: recreational basketball, competitive soccer) |
| Are there any sports you would like to play but avoid because of your knee? | Yes, No |
| Why do you avoid the activity? | Personal choice, Outside influence (parent, friend, coach, therapist, physician, etc.), Knee does not tolerate sport, Other (specify below) |
| Do you notice any stiffness or loss of motion in your knee? | Yes, No |
| How would you rate your pain on a scale of 0-10 at rest? (0 = No pain and 10 = worst pain imaginable) | 0, 1, 2, 3, 4, 5, 6, 7, 8 |
| How would you rate your pain on a scale of 0-10 during daily activities? (0 = No pain and 10 = worst pain imaginable) | 0, 1, 2, 3, 4, 5, 6, 7, 8 |
| How would you rate your pain on a scale of 0-10 during sport activities? (0 = No pain and 10 = worst pain imaginable) | 0, 1, 2, 3, 4, 5, 6, 7, 8 |

*This survey was utilized for several studies. Only patients of a single surgeon (SKA) who met inclusion and exclusion criteria were considered in the present study.

†If affirmative, confirmed via chart review.