Association Between Early Postoperative Graft Signal Intensity and Residual Knee Laxity After Anterior Cruciate Ligament Reconstruction

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**Background:** Magnetic resonance imaging (MRI) graft signal intensity is associated with graft damage after anterior cruciate ligament reconstruction (ACLR). However, little is known about the relationship between graft signal intensity and residual laxity of the reconstructed knee based on patient age.

**Purpose/Hypothesis:** To evaluate the relationship between graft signal intensity and residual laxity in younger and older patients who underwent ACLR. We hypothesized that higher graft signal intensity would be associated with reduced postoperative knee stability.

**Study Design:** Cohort study; Level of evidence, 3.

**Methods:** A total of 192 patients who underwent double-bundle ACLR were recruited. Proton density–weighted and T2-weighted MRI was performed at 3, 6, and 12 months after surgery, and the signal intensity ratio (SIR) of the anteromedial and posterolateral bundles was measured as the graft signal intensity reference values. At 12 months after surgery, if the KT-1000 arthrometer measurement exhibited a side-to-side difference of ≥2 mm, the patient was determined as having anterior knee laxity. Rotatory knee laxity was defined as a positive pivot shift with International Knee Documentation Committee grade ≥1. The Mann-Whitney U test was used to compare the SIR in patients with and without residual laxity. The Spearman correlation coefficient was used to evaluate the relationship between demographic parameters and the SIR. Based on receiver operating characteristic curves, the optimal SIR cutoff values to predict residual laxity were calculated, and logistic regression analysis was conducted.

**Results:** Of 192 patients, 26 (13.5%) had anterior knee laxity, and 20 (10.4%) had rotatory knee laxity. The SIR was negatively correlated with age. In younger patients (<30 years; n = 135), those with residual laxity had a significantly higher SIR than those without laxity; this relationship was not significant in older patients (≥30 years; n = 57). Based on receiver operating characteristic curves and logistic regression analysis, the cutoff values that were determined for the SIR were significantly associated with a higher odds ratio of residual laxity.

**Conclusion:** Graft signal intensity decreased with patient age. Patients with higher graft signal intensity in the early postoperative phase after ACLR exhibited a higher prevalence of residual laxity, particularly in younger patients.

**Keywords:** anterior cruciate ligament; reconstruction surgery; graft signal intensity; age; residual knee laxity
biomarkers such as blood samples and imaging studies would be useful to determine the risk of postoperative graft failure or residual knee laxity after anatomic ACLR as early as possible.

Graft signal intensity on magnetic resonance imaging (MRI) has been demonstrated to correspond to the progressive remodeling period of the ACL graft. Histologically, high graft signal intensity indicates hypervascularity or hypercellular tissue. This histologically higher graft signal can be regularly observed during the early postoperative phase (6 months after surgery) and gradually decreases between the first and second years after ACLR. Accordingly, this transition of graft signal intensity suggests a progressive graft remodeling period. In addition to the concept of the graft remodeling process, previous studies reported that the ACL graft signal increased as a result of damage to the reconstructed graft, caused by an acute graft-bending angle, tunnel malpositioning, and femoral notch impingement. Therefore, persistently high graft signal intensity might indicate graft deterioration, which could lead to worse clinical outcomes after ACLR. However, there is insufficient evidence to explain how higher graft signal intensity affects the clinical outcomes of patients who undergo anatomic ACLR. In particular, limited MRI graft signal data within the first year after ACLR are available regarding whether the graft signal affects future clinical outcomes.

The purpose of the current study was to evaluate the association between graft signal intensity during the first year after anatomic ACLR and the prevalence of residual knee laxity. Our hypothesis was that higher graft signal intensity would be associated with a greater prevalence of residual knee laxity at 12 months after ACLR.

METHODS

A total of 459 patients underwent anatomic double-bundle ACLR using a hamstring tendon autograft at our hospital between 2010 and 2018. We excluded patients with incomplete data, open physes, infections, multiple ligament injuries that required repair or reconstruction, and knee osteoarthritis defined as Kellgren-Lawrence grade ≥2 as well as those who underwent meniscal resection. In addition, patients who reinjured their ACL graft were also excluded. Ultimately, 192 patients were enrolled in the current retrospective cohort study. The main reason for exclusion was incomplete MRI data as the result of missing examinations. The study protocol was approved by the ethics committee of our institution, and all participants provided informed consent.

Surgical Procedure of Anatomic Double-Bundle ACLR

Anatomic double-bundle ACLR for the study participants was performed according to a previous study. The harvested semitendinosus tendon was split, and the distal and proximal halves of the tendon were looped and used as the anteromedial bundle (AMB) and posterolateral bundle (PLB), respectively. Each graft was 4.5 to 7.0 mm in diameter. Femoral tunnels for both the AMB and PLB were aimed posterior to the resident's ridge onto the direct insertion of the ACL using the transportal technique. A tibial tunnel was created as anteriorly as possible and positioned close to the bony ridge, which corresponded to the anterior boundary of the tibial ACL attachment (Parsons knob).

The proximal end of each graft was fixed by a fixed-loop cortical suspensory system (Position Suture Plate; B. Braun Aesculap) until August 2014; thereafter, an adjustable-loop cortical suspensory system (TightRope; Arthrex) was applied to retain the graft and provide a longer portion of the graft within the femoral tunnel to maximize the graft-bone interface area. The distal end of each graft was fixed by a suture mini-disc (B. Braun Aesculap) with maximum manual force applied as initial tension at 15° to 20° of flexion. Menisci were treated before double-bundle ACLR. An unstable meniscal injury in the red-red or red-white zone by probing was repaired as much as possible.

Postoperative rehabilitation was also consistent with a previous study. Briefly, crutch-assisted weightbearing ambulation and range of motion were permitted the day after ACLR, and if the patients underwent concomitant meniscal repair, their knee joint was immobilized in a brace for 1 week. Running, open kinetic chain exercises, and jump-landing training were allowed at 3 months; the patients started to return to sports activities at 6 to 9 months after surgery.

Clinical Assessment

At the 12-month follow-up, several surgeons at our sports medicine clinic examined all participants. All examiners regularly perform ACLR and thus have experience with this procedure, and the postoperative examination was conducted at a single institute. Sagittal knee stability was evaluated using a KT-1000 arthrometer with the Lachman test; if the KT-1000 arthrometer side-to-side
difference exhibited ≥2-mm anterior tibial translation relative to the femur, the knee was defined as having residual anterior knee laxity (AKL). For rotatory knee laxity (RKL), the pivot-shift test was administered, and it was assessed by International Knee Documentation Committee (IKDC) grading\textsuperscript{15,21}; if the patients exhibited an IKDC grade ≥1, they were defined as having residual RKL.

Measurement of Graft Signal Intensity on MRI

All patients underwent 1.5-T MRI of their operated knee joint at a mean of 3, 6, and 12 months after ACLR (Signa HDxt; GE Healthcare). The knee joint was kept immobilized, flexed at 10°, and externally rotated at 10° to 15° with a customized frame during the examination. Sagittal fast spin echo proton density–weighted (PDW) (repetition time [TR]/echo time [TE], 3000/98 milliseconds; slice thickness/intersection gap, 4.0/1.0 mm; field of view [FOV], 16 cm; number of excitations [NEX], 2; echo train length [ETL], 14; band width, ±31.25 kHz/FOV) and fast spin echo T2-weighted (T2W) (TR/TE, 3000/16 milliseconds; slice thickness/intersection gap, 4.0/1.0 mm; FOV, 16 cm; NEX, 2; ETL, 14; band width, ±31.25 kHz/FOV) sequences were obtained. The oblique sagittal image was reconstructed by defining the plane that passed through the centers of both the femoral and tibial tunnel outlets.

Based on the oblique sagittal image, signal intensity was measured by calculating the mean region of interest in the femoral and tibial tunnel outlets. The boundary of the measured region was determined by calculating the mean region of interest in the femoral and tibial tunnel outlets. The oblique sagittal image was reconstructed by defining the plane that passed through the centers of both the femoral and tibial tunnel outlets.

To further study the effect of age on the SIR, we divided the participants into younger (≤30 years; n = 135) and older (>30 years; n = 57) groups; 30 years was chosen because the histogram of participant age showed 2 peaks divided by this age (Appendix Figure A2). In the younger group, the SIR of patients with AKL (n = 19) was significantly higher than those without AKL (n = 116) at 3, 6, and 12 months after ACLR (Figure 1); the SIR of patients with RKL (n = 15) was significantly higher than those without RKL (n = 120) at 3 and 6 months after ACLR, whereas this relationship lost significance at 12 months after surgery (Figure 2). In the older group, the SIR of those without AKL (n = 50) and RKL (n = 52) was not associated with those having AKL (n = 7) and RKL (n = 5), respectively (Figures 1 and 2).

In the younger group, the ROC curve analysis indicated that the SIR within the first postoperative year was able to discriminate between patients with and without AKL at 12 months postoperatively (area under the curve, 0.676-0.848). In addition, the SIR at 3 and 6 months postoperatively was able to discriminate between patients with and without RKL (area under the curve, 0.672-0.763) (Table 3).

The optimal SIR cutoff values were determined based on the ROC curve analysis. Logistic regression analysis demonstrated that these optimal cutoff values were significantly associated with higher odds ratios of residual knee laxity only in the younger group (Table 4). For instance, in patients aged <30 years, the cutoff values of the SIR on PDW MRI at 3 months after ACLR demonstrated the highest odds (odds ratio, 17.5 [determined by AMB] or 23.1 [determined by PLB]) of a patient’s having AKL. According to the 3-month SIR of the AMB on PDW MRI, AKL was observed in 14 of 30 patients (46.7%) with an SIR of ≥1.25; on the other hand, AKL was observed in only 5 of 105 patients (4.8%) with an SIR of <1.25 (Table 4). Consistently, according to the 3-month SIR of the PLB on PDW MRI, AKL was observed in 12 of 20 patients (60.0%) with an SIR of ≥1.50; on the other hand, AKL was observed in 7 of 115 patients (6.1%) with an SIR of <1.50 (Table 4).
DISCUSSION

The primary finding of this study was that higher ACL graft signal intensity on MRI in the early postoperative phase (i.e., within the first year) was significantly associated with a higher prevalence of residual knee laxity at 1 year after anatomic double-bundle ACLR. Another important finding was that

| TABLE 1 |
| --- |
| Characteristics of Study Participants<sup>a</sup> |

|               | Anterior Knee Laxity Without (n = 166) | With (n = 26) | Rotatory Knee Laxity Without (n = 172) | With (n = 20) |
|---------------|----------------------------------------|--------------|----------------------------------------|--------------|
| Age, y        | 24.4 ± 11.4                            | 23.3 ± 10.9  | 24.6 ± 11.6                            | 21.5 ± 8.9   |
| Sex, male/female, n | 55/111                                 | 9/17         | 58/114                                 | 6/14         |
| Body mass index, kg/m² | 23.0 ± 3.5                             | 23.0 ± 4.0   | 23.1 ± 3.5                             | 22.6 ± 3.8   |
| Tegner score | 6.4 ± 1.6                              | 6.5 ± 2.1    | 6.4 ± 1.7                              | 6.5 ± 1.9    |
| Time from injury to surgery, mo | 5.5 ± 23.1                          | 3.5 ± 4.7    | 5.5 ± 22.7                             | 2.9 ± 2.9    |
| KT-1000 arthrometer side-to-side difference, mm | 0.5 ± 0.5                           | 2.2 ± 0.4<sup>b</sup> | 0.6 ± 0.6                             | 1.8 ± 0.9<sup>b</sup> |

**Signal intensity ratio**

|               | AMB on PDW |                         |                         |
|---------------|------------|-------------------------|-------------------------|
| 3 mo          | 1.25 ± 0.45<sup>b</sup> | 1.00 ± 0.27             | 1.24 ± 0.42<sup>b</sup> |
| 6 mo          | 1.54 ± 0.50<sup>b</sup> | 1.23 ± 0.33             | 1.53 ± 0.54<sup>b</sup> |
| 12 mo         | 1.36 ± 0.32<sup>b</sup> | 1.22 ± 0.40             | 1.31 ± 0.35             |

|               | AMB on T2W |                         |                         |
|---------------|------------|-------------------------|-------------------------|
| 3 mo          | 1.22 ± 0.28<sup>b</sup> | 1.06 ± 0.19             | 1.19 ± 0.30             |
| 6 mo          | 1.33 ± 0.25<sup>b</sup> | 1.15 ± 0.23             | 1.32 ± 0.25<sup>b</sup> |
| 12 mo         | 1.14 ± 0.22 | 1.08 ± 0.24             | 1.11 ± 0.22             |

|               | PLB on PDW |                         |                         |
|---------------|------------|-------------------------|-------------------------|
| 3 mo          | 1.38 ± 0.52<sup>b</sup> | 1.08 ± 0.30             | 1.38 ± 0.47<sup>b</sup> |
| 6 mo          | 1.70 ± 0.48<sup>b</sup> | 1.33 ± 0.33             | 1.65 ± 0.53<sup>b</sup> |
| 12 mo         | 1.56 ± 0.38<sup>b</sup> | 1.32 ± 0.40             | 1.50 ± 0.43             |

|               | PLB on T2W |                         |                         |
|---------------|------------|-------------------------|-------------------------|
| 3 mo          | 1.32 ± 0.36<sup>b</sup> | 1.11 ± 0.25             | 1.27 ± 0.33<sup>b</sup> |
| 6 mo          | 1.53 ± 0.44<sup>b</sup> | 1.23 ± 0.29             | 1.46 ± 0.46<sup>b</sup> |
| 12 mo         | 1.31 ± 0.33<sup>b</sup> | 1.16 ± 0.30             | 1.26 ± 0.36             |

<sup>a</sup>Data are reported as mean ± SD unless otherwise indicated. AMB, anteromedial bundle; PDW, proton density weighted; PLB, posterolateral bundle; T2W, T2 weighted.

<sup>b</sup>Statistically significant difference between patients with and without laxity (.05; Mann-Whitney U test).

| TABLE 2 |
| --- |
| Spearman Correlation Coefficients for Association Between Patient Characteristics and Signal Intensity Ratio<sup>a</sup> |

|               | r Value |
|---------------|---------|
| Age           |         |
| Body Mass Index |       |
| Tegner Score  |         |
| Time From Injury to Surgery |       |

|               | AMB on PDW |                         |                         |
|---------------|------------|-------------------------|-------------------------|
| 3 mo          | -0.296<sup>b</sup> | -0.010                  | 0.217<sup>b</sup>       | 0.079 |
| 6 mo          | -0.287<sup>b</sup> | 0.051                   | 0.220<sup>b</sup>       | 0.007 |
| 12 mo         | -0.142<sup>b</sup> | 0.132                   | 0.138                  | <0.001 |

|               | AMB on T2W |                         |                         |
|---------------|------------|-------------------------|-------------------------|
| 3 mo          | -0.249<sup>b</sup> | -0.038                  | 0.121                  | 0.151<sup>b</sup> |
| 6 mo          | -0.211<sup>b</sup> | 0.067                   | 0.222<sup>b</sup>       | -0.043 |
| 12 mo         | -0.033      | 0.102                   | 0.051                  | 0.069 |

|               | PLB on PDW |                         |                         |
|---------------|------------|-------------------------|-------------------------|
| 3 mo          | -0.329<sup>b</sup> | -0.044                  | 0.209<sup>b</sup>       | 0.060 |
| 6 mo          | -0.357<sup>b</sup> | -0.024                  | 0.240<sup>b</sup>       | 0.023 |
| 12 mo         | -0.168<sup>b</sup> | 0.065                   | 0.150<sup>b</sup>       | -0.001 |

|               | PLB on T2W |                         |                         |
|---------------|------------|-------------------------|-------------------------|
| 3 mo          | -0.307<sup>b</sup> | -0.046                  | 0.160<sup>b</sup>       | 0.134 |
| 6 mo          | -0.214<sup>b</sup> | -0.018                  | 0.194<sup>b</sup>       | 0.006 |
| 12 mo         | -0.051      | 0.025                   | 0.022                  | 0.050 |

<sup>a</sup>Based on the r value, the correlations were interpreted as poor to fair. AMB, anteromedial bundle; PDW, proton density weighted; PLB, posterolateral bundle; T2W, T2 weighted.

<sup>b</sup>Statistically significant (.05).

**DISCUSSION**

The primary finding of this study was that higher ACL graft signal intensity on MRI in the early postoperative phase (i.e., within the first year) was significantly associated with a higher prevalence of residual knee laxity at 1 year after anatomic double-bundle ACLR. Another important finding was that
Graft signal intensity was negatively correlated with the patient's age. For the patients younger than 30 years, those with higher graft signal intensity demonstrated a significantly higher prevalence of residual knee laxity. Conversely, in older patients (≥30 years), higher graft signal intensity was not significantly associated with residual knee laxity.

Figure 1. Comparison of graft signal intensity between participants with anterior knee laxity (AKL+) and without anterior knee laxity (AKL−) based on ligament bundle, age group, and imaging sequence. *Statistically significant difference between patients with and without laxity (P < .05; Mann-Whitney U test). AMB, anteromedial bundle; PDW, proton density–weighted; PLB, posterolateral bundle; T2W, T2-weighted.
Of the patients who underwent anatomic double-bundle ACLR using a hamstring tendon autograft, those with higher graft signal intensity in the early postoperative phase, such as at 3 or 6 months, showed a significantly higher prevalence of residual knee laxity at 1 year after surgery. With regard to the previous data, Hakozaki et al.\textsuperscript{14} reported that, using a similar outcome of residual knee laxity as that of the current study, higher graft signals

Figure 2. Comparison of graft signal intensity between participants with rotatory knee laxity (RKL\textsuperscript{+}) and without rotatory knee laxity (RKL\textsuperscript{−}) based on ligament bundle, age group, and imaging sequence. *Statistically significant difference between patients with and without laxity (P < .05; Mann-Whitney U test). AMB, anteromedial bundle; PDW, proton density–weighted; PLB, posterolateral bundle; T2W, T2-weighted.
### TABLE 3
Results of Receiver Operating Characteristic Curve Analysis of Signal Intensity Ratio and Residual Knee Laxity at 1 Year Postoperatively

| Age <30 y (n = 19) | Age ≥30 y (n = 7) | Age <30 y (n = 15) | Age ≥30 y (n = 5) |
|---------------------|-------------------|-------------------|-------------------|
| **Anterior Knee Laxity** | **Rotatory Knee Laxity** | **Anterior Knee Laxity** | **Rotatory Knee Laxity** |
| AMB on PDW | | | |
| 3 mo | 0.811 | 0.411 | 0.718 | 0.638 |
| (0.691-0.932)<sup>a</sup> | (0.177-0.645) | (0.542-0.893)<sup>a</sup> | (0.403-0.974) |
| 6 mo | 0.785 | 0.420 | 0.688 | 0.558 |
| (0.680-0.891)<sup>a</sup> | (0.194-0.646) | (0.527-0.850)<sup>a</sup> | (0.317-0.799) |
| 12 mo | 0.676 | 0.529 | 0.611 | 0.558 |
| (0.561-0.791)<sup>a</sup> | (0.327-0.730) | (0.465-0.756) | (0.315-0.800) |
| AMB on T2W | | | | |
| 3 mo | 0.791 | 0.497 | 0.695 | 0.477 |
| (0.679-0.904)<sup>a</sup> | (0.296-0.699) | (0.522-0.868)<sup>a</sup> | (0.232-0.722) |
| 6 mo | 0.779 | 0.597 | 0.763 | 0.592 |
| (0.671-0.888)<sup>a</sup> | (0.397-0.798) | (0.633-0.893)<sup>a</sup> | (0.353-0.832) |
| 12 mo | 0.613 (0.479-0.748) | 0.563 | 0.581 | 0.535 |
| (0.324-0.801) | (0.415-0.746) | (0.323-0.746) | |
| PLB on PDW | | | | |
| 3 mo | 0.833 | 0.374 | 0.718 | 0.723 |
| (0.711-0.955)<sup>a</sup> | (0.144-0.605) | (0.548-0.889)<sup>a</sup> | (0.523-0.923) |
| 6 mo | 0.848 | 0.509 | 0.734 | 0.485 |
| (0.758-0.939)<sup>a</sup> | (0.276-0.741) | (0.587-0.881)<sup>a</sup> | (0.220-0.750) |
| 12 mo | 0.747 | 0.526 | 0.654 | 0.588 |
| (0.642-0.853)<sup>a</sup> | (0.287-0.764) | (0.498-0.810) | (0.371-0.806) |
| PLB on T2W | | | | |
| 3 mo | 0.804 | 0.460 | 0.672 | 0.554 |
| (0.688-0.919)<sup>a</sup> | (0.244-0.676) | (0.495-0.849)<sup>a</sup> | (0.288-0.820) |
| 6 mo | 0.847 | 0.534 | 0.711 | 0.681 |
| (0.771-0.923)<sup>a</sup> | (0.292-0.777) | (0.577-0.845)<sup>a</sup> | (0.506-0.856) |
| 12 mo | 0.686 | 0.577 | 0.619 | 0.581 |
| (0.557-0.814)<sup>a</sup> | (0.343-0.812) | (0.448-0.791) | (0.422-0.739) |

<sup>a</sup>Data are reported as area under the curve (95% CI). AMB, anteromedial bundle; PDW, proton density weighted; PLB, posterolateral bundle; T2W, T2 weighted.

<sup>b</sup>Statistically significant for discriminating between patients with and without residual knee laxity at 12 months postoperatively (P < .05).

### TABLE 4
Results of Logistic Regression Analysis for Signal Intensity Ratio Cutoff Values and ORs for Residual Knee Laxity in Patients Aged <30 Years

| Cutoff B | OR (95% CI) | Cutoff B | OR (95% CI) |
|---------|------------|---------|------------|
| **Anterior Knee Laxity** | **Rotatory Knee Laxity** | **Anterior Knee Laxity** | **Rotatory Knee Laxity** |
| AMB on PDW | | | |
| 3 mo | 1.25 | 2.862 | 17.5 (5.5-55.2) | 1.25 | 2.303 | 10.0 (3.1-32.4) |
| 6 mo | 1.65 | 2.012 | 7.5 (2.6-21.4) | 1.65 | 1.601 | 5.0 (1.6-15.4) |
| 12 mo | 1.30 | 1.302 | 3.7 (1.3-10.4) | NS | NS | |
| AMB on T2W | | | | |
| 3 mo | 1.10 | 2.453 | 11.6 (2.6-52.7) | 1.13 | 1.888 | 6.6 (2.0-22.0) |
| 6 mo | 1.25 | 2.120 | 8.3 (2.6-26.9) | 1.20 | 2.277 | 9.8 (2.1-45.2) |
| 12 mo | NS | NS | | NS | NS | |
| PLB on PDW | | | | |
| 3 mo | 1.50 | 3.142 | 23.1 (7.1-75.1) | 1.50 | 2.331 | 10.3 (3.2-33.4) |
| 6 mo | 1.85 | 2.921 | 18.6 (5.8-59.2) | 1.55 | 1.820 | 6.2 (1.8-20.7) |
| 12 mo | 1.35 | 2.488 | 12.0 (2.7-54.6) | 1.55 | 1.333 | 3.8 (1.3-11.5) |
| PLB on T2W | | | | |
| 3 mo | 1.35 | 2.724 | 15.2 (4.9-47.6) | 1.35 | 2.187 | 8.9 (2.8-28.7) |
| 6 mo | 1.50 | 2.446 | 11.5 (3.9-33.9) | 1.25 | 1.905 | 6.7 (1.5-31.1) |
| 12 mo | 1.35 | 1.292 | 3.6 (1.3-10.0) | 1.35 | 1.201 | 3.3 (1.1-10.1) |

<sup>a</sup>AMB, anteromedial bundle; NS, not significant; OR, odds ratio; PDW, proton density weighted; PLB, posterolateral bundle; T2W, T2 weighted. Cutoff value indicates the signal intensity ratio.
on PDW and T2W MRI at 1 year after anatomic double-bundle ACLR were associated with residual AKL defined by a ≥4-mm side-to-side difference using the KT-2000 arthrometer. Additionally, a more recent clinical study reported by Putnis et al32 consistently demonstrated that patients with higher graft signal intensity around the femoral tunnel aperture exhibited a higher risk of graft reruptures after single-bundle ACLR. The current study significantly supports these previous data in which higher graft signal intensity is associated with greater laxity derived from intrinsic damage to the reconstructed ACL graft after reconstruction.1,8,18,25,32,33 Therefore, knee surgeons should pay specific attention to the risk of not only graft reruptures but also residual knee laxity in patients who exhibit higher graft signal intensity on MRI during the first postoperative year.

Moreover, regarding the clinical relevance of ACL graft signal intensity, the current data provide insight on 2 important aspects that should be considered. First, the ability of 3- or 6-month postoperative SIRs to detect residual knee laxity was superior to that of 12-month SIRs (Tables 3 and 4). At this point, few previously published studies support the current finding that earlier postoperative graft signal intensity (ie, within 6 months after ACLR) is superior in detecting future residual knee laxity. It should be noted that several MRI-based studies have reported that the mean graft signal intensity at 3 months after ACLR was lower than that at 6 months because the tissue reparative and revascularization processes are in progress. Thereafter, the mean graft signal intensity is likely to peak around 6 months after reconstruction and then gradually decline and return to near normal levels at 2 years after surgery, corresponding to the healing process of the reconstructed graft.29,30,46 Based on the initial process of MRI graft signal intensity, an earlier postoperative period such as 3 months is less likely to be affected by the healing process of the graft. Therefore, a higher graft signal in the early postoperative period can be a more specific and appropriate finding to indicate graft deterioration, which may predict residual knee laxity.

Another important finding is that patient age was associated with graft signal intensity; graft signal intensity of older patients was lower than that of younger patients. Notably, a recent study by Putnis et al32 demonstrated that patient age was negatively correlated with graft signal intensity on MRI. Considering the current and previous data, older patients are less likely to experience similar revascularity bioactivity or tissue reparative reactions compared with younger patients.29,30,33,46 Moreover, older patients are generally less active and less likely to stress their reconstructed ACL graft compared with younger patients. Knee surgeons should therefore consider patient age when evaluating graft signal intensity after ACLR; older patients attenuate the pathological significance of graft signal intensity with respect to predicting residual knee laxity.

The current data indicated the clinical relevance of postoperative MRI graft signal intensity that could be useful to validate the prognosis of patients who undergo anatomic ACLR. In particular, younger patients with higher physical activity levels would benefit from undergoing follow-up MRI between 3 and 6 months after ACLR to determine the risk of developing residual knee laxity. If the graft signal is far higher than its mean value, surgeons should pay attention to the risk of residual knee laxity and alter postoperative treatment to preserve the graft, for instance, by delaying the progression of rehabilitation. Future MRI studies are required to further explain the mechanisms underlying graft signal intensity after anatomic ACLR.

As for the concept of double-bundle ACLR, previous biomechanical studies have put an emphasis on the different functions of the AMB and PLB in restraining anterior laxity and controlling tibial axial rotation laxity.2,13,34,36 The AMB has been reported to have a more specific biomechanical behavior to stabilize anterior tibial translation with the knee flexed past 30°.2,36 On the other hand, the PLB has been reported to stabilize anterior tibial translation when the knee is at full extension or 15° of flexion.2,36 and the oblique orientation of the PLB in the coronal plane allows greater ability to stabilize tibial rotatory laxity than a more vertical AMB.2,13,34 Notably, the current data conflict with these previous studies. In other words, the SIRs of the AMB could predict the development of both AKL and RKL; consistently, that of the PLB could predict the development of both AKL and RKL. However, previous studies have reported that both the AMB and PLB are important for stabilizing anterior and rotatory tibial laxity relative to the femur after ACLR. In fact, 2 clinical studies using navigation systems clarified that isolated fixation of either the AMB or PLB reduced anterior tibial translation against both anterior tibial loading and tibial internal rotation at the same time.22,24 The findings of the current clinical study support the latter 2 navigation system studies22,24; for patients who undergo ACLR, both the AMB and the PLB have an important role to stabilize AKL and RKL.

**Limitations**

The current study had several limitations. First, the graft that was harvested in the current study was a hamstring tendon autograft. Therefore, the current findings would not be applicable to ACL-injured patients reconstructed with an allograft or bone–patellar tendon–bone graft. Second, the follow-up period was relatively short, up to 12 months after surgery, and further relationships remain unclear. In addition, we did not include laxity data at 3 and 6 months after ACLR. Third, RKL was evaluated using IKDC grading. Quantitative methods16,26 would clarify the detailed mechanism between RKL and graft signal intensity. Fourth, the current study did not obtain an enhanced MRI sequence, which would be more effective for evaluating the revascularization or tissue reparative process.10,29,30 Fifth, we switched from a fixed-loop cortical suspension device to an adjustable-loop device to fix the proximal end of the hamstring tendon graft. This may have affected the process of graft healing and its following alteration of the graft signal on MRI. Finally, multiple factors are associated with the development of residual laxity after ACLR, the most important being nonanatomic tunnel placement.5,19,48 Other factors include a concomitant meniscal tear with the
ACL injury as well as intrinsic joint laxity before the ACL injury. Future large-sample cohort studies that include graft signal intensity data would reveal the interrelationship among these multiple factors in the development of residual laxity after ACLR.

CONCLUSION

The current study elucidated the relationship between graft signal intensity on MRI and residual knee laxity after anatomic double-bundle ACLR. The graft signal was negatively correlated with patient age. In patients younger than 30 years, higher graft signal intensity in the early postoperative period (between 3 and 6 months after ACLR) was associated with a higher prevalence of residual knee laxity at 1 year postoperatively.

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APPENDIX

Appendix Figure A1. Example of evaluating graft signal intensity on magnetic resonance imaging. The signal intensity ratio was calculated as (AMB intensity or PLB intensity)/PCL intensity. AMB, anteromedial bundle; PCL, posterior cruciate ligament; PLB, posterolateral bundle.

Appendix Figure A2. Histogram of patient age.