Simultaneous measurements of sagittal knee laxity with an external device and radiostereometric analysis
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We obtained simultaneous measurements of sagittal knee laxity in 12 consecutive patients after reconstruction of the anterior cruciate ligament (ACL), using the Stryker laxity tester and radiostereometric analysis (RSA).

The mean anteroposterior (AP) displacement when a 90 N load was applied in both directions was 5.3 ± 2.7 mm with RSA and 9.8 ± 1.6 mm with the external device (p < 0.001). The corresponding measurements at a load of 180 N were 5.7 ± 2.4 mm and 13.8 ± 3.7 mm, respectively (p < 0.001).

More than 50% of the sagittal knee movement, as measured by the external device at a load of 180 N, was not true femorotibial displacement of the joint but was due to soft-tissue deformation.

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Commercial devices for measuring AP laxity of the knee have been shown to be inaccurate.1-3 Soft-tissue deformation has been suggested as the reason for this error.4,5 Our aim was to evaluate the accuracy of one such device and assess the magnitude of the error during measurements in vivo. We measured the anteroposterior (AP) displacement with this appliance, and compared it with simultaneous measurement using radiostereometric analysis (RSA).6

Patients and Methods

We studied 12 consecutive patients three years after reconstruction of an anterior cruciate ligament. There were four women and eight men with a mean age at surgery of 25 years (17 to 37). The operations had been performed through a miniarthrotomy using the central third of the patellar tendon. Four tantalum markers had been implanted in the distal femur and four in the proximal tibia. The study was approved by the Ethics Committee of Lund University Hospital.

We performed simultaneous measurements of the total AP laxity at the three-year postoperative follow-up using the Stryker system (Orthopedic System Inc, Hayward, California) and RSA. Simultaneous RSA-radiographs were obtained in the frontal and lateral projections while applying a posterior force of 90 N and 180 N as measured on the Stryker laxity tester. Radiographs were also obtained while applying an anterior force of 90 N and 180 N. All measurements were made with the knee in 25 ± 5° of flexion. The marker images on all radiographs were digitised using a precision digitising table (Hasselblad Engineering Inc, Göteborg, Sweden). Movement of the tibia relative to the femur was obtained by the KINEMA routine as described by Fridén, Ryd and Lindstrnad.7 We calculated the total AP displacement by each method at loads of both 90 N and 180 N. The actual knee angles after application of the different loads were also obtained from the RSA calculations.

The accuracy of the RSA technique has previously been determined as ±0.2 mm.8 The precision, including biological variation (different muscle relaxation, positioning, etc), has also been calculated as ± 2SD, and found to be ±1.6 mm9 and ±2.2 mm.7 For the Stryker device the precision was found to be less than 3.2 mm.3

Statistical analysis used a paired Student’s t-test and the relationship between the results from the two testing loads and methods was expressed by Pearson’s correlation coefficient (r). The 95% confidence interval (CI) for the mean differences between the methods and the 95% limits of agreement were calculated according to the method of Bland and Altman.10 The results are given as mean values ±SD and p values of less than 0.05 (two-tailed tests) were considered significant.

Results

There was no increase in measured AP movement between the RSA measurements at 90 N and 180 N (5.3 ± 2.7 mm at
90 N and 5.7 ± 2.4 mm at 180 N). Simultaneous recordings using the external device showed increasing laxity with increasing load: 9.8 ± 1.6 mm at 90 N and 13.8 ± 3.7 mm at 180 N (p < 0.001; Table I, Fig. 1). The correlation coefficient (r) was high within each of the two methods when 90 N and 180 N loads were compared: RSA 0.92 (p < 0.01), Stryker 0.83 (p < 0.01). Between the two methods, however, no significant correlation coefficients were found: at 90 N r = 0.30 and at 180 N r = 0.20. The mean difference between the two methods was 4.4 mm at 90 N and 8.0 mm at 180 N.

There was no clear evidence of a relationship between the individual difference between the methods (bias = systematic error) and the individual mean total AP displacement; at 90 N r = –0.50 and at 180 N r = 0.47 (NS, Figs 2 and 3). The 95% CI for the bias was 3 to 6 mm at 90 N and 5 to 11 mm.

Table I. Total AP displacement (mm) and change in flexion angle (degrees) in 12 patients before surgery and three years after ACL reconstruction as measured by RSA and the Stryker laxity tester

<table>
<thead>
<tr>
<th>Case</th>
<th>RSA</th>
<th>Stryker</th>
<th>RSA</th>
<th>Stryker</th>
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<th>180N postop</th>
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Fig. 1
Total mean (±SD) AP displacement at loads of 90 N and 180 N three years after ACL reconstruction measured simultaneously by RSA and the Stryker laxity tester.

Fig. 2
Comparison of RSA and the Stryker laxity tester according to Bland and Altman at two different loads, 90 N and 180 N, respectively (— 95% CI for the mean differences between the methods; ... 95% limits of agreement).
at 180 N. The corresponding 95% limits of agreement were -1 to +10 mm at 90 N, and 0 to 16 mm at 180 N.

We found a force-induced change in knee flexion between anterior and posterior loads as measured by RSA; 8.4 ± 2.7° at 90 N and 13.3 ± 3.9° at 180 N (Table I). When this angular change was related to each individual laxity difference between RSA and the external device, \( r = 0.51 \) (NS) at 90 N and \( r = 0.62 \) at 180 N (NS).

**Discussion**

The evaluation of different techniques for laxity measurements has mainly focused on the reproducibility of the methods. Only a few studies have determined the accuracy. Using simultaneous measurements by RSA and an external device on two specimens in vitro Edixhoven et al.\(^{11}\) found smaller displacements on RSA. In 36 patients with torn anterior cruciate ligaments, Kärholm et al.\(^{9}\) also found smaller displacements with RSA than had been earlier described for external devices.\(^{12,13}\) Jonsson, Kärholm and Emqvist\(^{14}\) compared RSA with the KT-1000 laxity tester (MEDMetric Corporation, San Diego, California) in 94 knees with chronic injury of the anterior cruciate ligament of which 55 had undergone ligament reconstruction. Contrary to our results, the reconstructed knees showed smaller displacements with the KT-1000 tester. The measurements were, however, not performed simultaneously and smaller loads were applied with the KT-1000 tester than for the RSA measurements; this made the comparison between the two methods inappropriate.

We have previously used RSA and measurements with external devices in a smaller group of patients.\(^{5}\) The measurements were not performed simultaneously and we showed significantly larger displacements with the Stryker laxity tester only when the greater force of 180 N was used. Shino et al.\(^{4}\) demonstrated in vitro that soft-tissue deformation was responsible for a large proportion of the laxity measured by an external device, and Steiner et al.\(^{3}\) found that the ‘testing error’ accounted for more than 50% of the measured difference in displacement between right and left knees of normal subjects. Radiological recordings of the displacement induced by passive forces have also been analysed,\(^{15-17}\) and the possible errors are the same as with external devices, except for the soft-tissue deformation. In addition, there are difficulties in identifying definite bony landmarks and compensating for magnification.\(^{17}\) To determine accurately the reliability of external devices in vivo, simultaneous measurements using RSA with implanted markers is probably the best technique. The error produced by intra-individual, repeated measurements was eliminated in the present study since the measurements were performed simultaneously, and thus errors, such as limb fixation,\(^{18}\) knee flexion,\(^{1,18}\) the degree of muscle relaxation,\(^{9,11}\) the starting position sagittally,\(^{5,9,11}\) and the day-to-day difference\(^{11,20}\) were excluded by the design of the study.

The finding of a low correlation between RSA and the external device agrees with earlier studies comparing radiological methods and such machines. Stäubli and Jakob\(^{21}\) examined 16 patients with clinically diagnosed ACL-deficient knees by simultaneous measurements with radiography and the KT-1000 arthrometer during epidural anaesthesia. Comparisons were made at 89 N, applied in the anterior direction. No correlation was found between the measurements obtained with the two methods.

The higher force (180 N) did not significantly increase the displacement as measured with RSA, and thus the RSA measurements verified a definite endpoint of joint displacement which could not be established using the external device in contrast to findings in most previous studies.\(^{1,3}\)

The difference between RSA and the external device became greater with increasing load, and at both 90 N and 180 N the device recorded significantly larger movement than the RSA measurements of the true skeletal displacement.

RSA is mainly suitable for scientific purposes owing to the need for implanted markers. It can, however, provide us with an important technique to assess the accuracy of different external measuring devices.

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**References**


