Congenital Absence of the Anterior Cruciate Ligament


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Author Affiliation | Disclosures

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Congenital absence of the anterior cruciate ligament (ACL) is a rare occurrence and has been seen most often in conjunction with conditions such as knee dislocation, knee dysplasia, proximal focal femoral deficiency, and fibular hemimelia.

We report on the incidental finding of ACL aplasia in a patient with a medial meniscal tear and history of leg-length discrepancy. Similar to earlier cases, this patient had hypertrophy of the meniscofemoral ligament of Humphrey, which likely provided stability. This case report emphasizes the importance of distinguishing between a stable and an unstable knee in congenital absence of the ACL. The patient provided written informed consent for print and electronic publication of this case report.

Case Report

A 20-year-old woman presented for orthopedic evaluation with worsening medial left knee pain. Her pain was intermittent in nature, occurring about every 1 to 2 months and of 1 to 2 days’ duration. Onset was while using the elliptical machine, walking on uneven ground, or navigating stairs. She denied any buckling, catching, locking, instability, or swelling.

Her history was significant for a breech delivery and leg anisomelia, for which she had a contralateral distal femoral and proximal tibial percutaneous epiphysiodesis performed at age 10 years. Family history was negative for limb deformities.

Physical examination was notable for absence of global ligamentous laxity, overall valgus alignment of the left lower extremity, minimally decreased motion, trace effusion, positive medial joint line tenderness, positive McMurray test, and 1+ Lachman test with guarding on pivot shift testing.

Plain films showed valgus alignment with narrowing of the lateral compartment, narrow intercondylar notch, and hypoplasia of the tibial eminences and lateral femoral condyle (Figure 1). Magnetic resonance imaging showed a large tear in the posterior horn of the medial meniscus, hypertrophy of the meniscofemoral ligament of Humphrey (Figure 2A), and nonvisualization of the ACL with a small remnant (Figure 2B).
Arthroscopy showed complete absence of fibers of the ACL, hypertrophy of the meniscofemoral ligament of Humphrey, and a large posterior horn medial meniscal tear. A partial medial meniscectomy was performed. More than 2 years after surgery, the patient was doing very well without pain or instability, and was exercising regularly without difficulty.

Discussion

Our patient had left-sided congenital absence of the ACL with associated limb-length discrepancy of more than 2.5 cm. Isolated absence of the ACL has been described in a few case reports in the literature. Congenital ACL absence has most often been found in association with conditions such as knee dislocation (occurring with a frequency of .017/1000 births),\(^1\) knee dysplasia,\(^2\)\(^,\)\(^3\) fibular hemimelia,\(^4\) and proximal focal femoral deficiency.\(^5\) Johansson and Aparisi\(^5\)\(^,\)\(^6\) linked the finding of ACL absence with instability in those patients with known limb-length discrepancy and symptomatic instability. This report presents a patient who has congenital absence of the ACL in a foreshortened limb and torn medial meniscus. The classification of the patient’s cruciate dysplasia would be type I, as described by Manner and colleagues.\(^7\) The incidence of meniscal tears in association with congenital ACL absence is unknown. There have been reports of absence of the ACL associated with a ring meniscus,\(^8\) absence of both cruciate ligaments and menisci,\(^9\) and a bucket-handle tear of the medial meniscus.\(^10\)

Gabos and colleagues\(^4\) recommend reconstructive surgery for patients with congenital absence of the ACL and
symptomatic knee instability. Limb lengthening/shortening and realignment procedures have allowed patients such as ours to have functionally anatomic limbs and high activity levels. Surgical treatment is pursued to restore mechanical alignment and stability. Our patient had no symptoms of instability.

Similar to 3 of the 4 patients presented by Gabos and colleagues, our patient had marked hypertrophy of the meniscofemoral ligament of Humphrey. The report by Gabos and colleagues of this finding was the first in the literature. The hypertrophy of this ligament suggests it has a role in stabilizing the knee with a congenitally absent ACL. Our patient had no instability in her left knee but presented because of episodes of pain.

Of significant concern is the long-term outcome of patients with congenital ACL aplasia. Crawford and colleagues reported 11 patients with ACL deficiency and fibular hemimelia at a mean age of 37 years, showing similar functional outcomes to age-matched controls. However, there was no radiographic follow-up reported in regard to the development of osteoarthritis. To our knowledge, there have been no series published comparing surgical and nonsurgical treatment of congenital absence of the ACL. In the study by Gabos and colleagues, all patients were treated with reconstruction because these patients had symptomatic instability.

**Conclusion**

This report presents a patient whose symptoms improved after resection of her medial meniscal tear. This patient will be followed long-term to delineate her clinical course and to monitor for instability and/or development of osteoarthritis. Future studies should compare the treatment of congenital absence of the ACL with reconstruction and with conservative management.

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