Etiology of the bifid (double) femoral head, with prior history of developmental dysplasia of the hip

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ABSTRACT

Our patient presented with a double femoral head, that is, two separate heads with individual epiphyses, but a single contiguous metaphysis. Two similar cases had been described in the literature. The only common feature in these three cases is that they had open reduction for Developmental Dysplasia of the Hip (DDH) through an anterior approach. No other pathology was detected in these patients. A rabbit model was created in which the cartilaginous anlage of the rabbit femoral head was surgically split. After 2 - 4 weeks a bifid femoral head developed, mimicking that described in the literature. We suggest that inadvertent damage to the femoral head during surgery for DDH may in fact lead to the development of a bifid femoral head. Prior history of DDH should be considered when the isolated bifid femoral head is identified.

Keywords: Bifid; Femoral Head; Etiology; Rabbit Study

1. INTRODUCTION

Bifid femoral heads are an unusual and puzzling deformity. While the etiology of the deformity is unknown, Ferguson [1], Rossman [2] and Salter [3] proposed that since increased pressure or forced positions contribute to the development of avascular necrosis of the hip, a tight iliopsoas tendon, or focal necrosis through forced positions, may be responsible for the development of a bifid femoral head. However, if a tight iliopsoas muscle is the major factor in producing a bifid femoral head then the condition should be common in the cerebral palsy population. Yet it has never been described in those patients. Variants of bifid femoral heads are described in the literature, such as seen in Perthe’s Disease and Meyer Dysplasia, and in a recent article by Osuji, et al. [4].

Two patients with bifid femoral heads, with prior history of surgical treatment for D.D.H. have been described in the literature [1,2]. Ours is the third similar case to be reported.

2. CASE REPORT

An 8-year-old girl presented for evaluation of left hip pain. She felt this at times when walking, and also on turning over in bed. Open reduction though an anterior approach was performed for D.D.H. as an infant. She walked with a mild limp, with the limb in neutral alignment, but ran rapidly with her limb internally rotated. Radiographs revealed a split femoral head with a larger anterior segment and a smaller posteromedial segment (Figures 1 and 2). The smaller femoral head was contained in the acetabulum with the hip in neutral alignment, and the larger head with the hip internally rotated. This explained why she felt pain when walking, as the smaller head was in the acetabulum. She internally rotated her limb when running and had no pain, as the larger head was in the acetabulum. Intraoperatively the iliopsoas tendon was not significantly tight, and the labrum was not deformed. The smaller head was excised. The larger head was reduced by internal rotation into its most optimal position in the acetabulum, and a derotational osteotomy was performed with internal fixation (Figure 3). She made excellent progress. She had mild residual shortening of the limb. She participated in sporting activities and at 8-year follow-up, she was a cheerleader.

3. OBJECTIVE

The common thread in our patient, and the two prior cases, is surgery at a very young age for developmental dysplasia of the hip. We propose that the deformity may be related to damage to the ossific nucleus of the femoral head during surgery for DDH. We therefore devised an
animal model to determine the effect of splitting the ossific nucleus, with a simple incision, on development of the femoral head.

4. METHODS

Approval for the following study was received from IRB.

Eight New Zealand White (NZW) rabbits underwent bisection of the right femoral ossific nucleus at 13 or 15 days of age. At this age, the 5 mm rabbit femoral head is visible and surgical technique is thus reproducible [5]. Through a medial approach the right cartilaginous femoral head was exposed and bisected under direct visualization, and the joint capsule and wound closed. The left hip was used as the control. Radiographs of both hips were obtained at 2 - 4 weeks intervals until the rabbits were euthanized at maturity, around 4 months. Both femoral heads and both respective acetabulae were resected and inspected grossly and histologically.

5. RESULTS

Two anesthetic deaths reduced the sample number to six rabbits. One had to be sacrificed before maturity because of illness. In three of the rabbits radiographs demonstrated double femoral heads, and in all three the secondary femoral head articulated with a pseudo-acetabulum (Figure 4). In the remaining three rabbits the operated-on femoral head was normal in one, demonstrated an elongated neck in the second and had an SCFE-like pattern in the third. The femoral heads with their respective acetabulae were dissected. A thick joint capsule was present around the bifid femoral heads, with a flexion contracture in the double-headed femurs, but no tight structures were visible.

Gross sections of each specimen revealed two truly bifid femoral heads (Figure 5), with each head lined with cartilage and separate epiphyses (Figure 6). The third double head, suspected by radiographic evidence, was determined to be only a spicule of bone without any cartilaginous surface.
Histology showed all controls to have hemispherical heads. Two bifid femoral heads were present (Figure 7). One of these specimens showed a couple of dead bone spicules at the cleft site, however this was due to remodeling and not consistent with osteonecrosis. Another specimen showed one head with a second head-like structure, which was actually composed of compact bone without a physis. The fourth specimen demonstrated an elongated epiphysis without any scar tissue. The fifth specimen had a widened acetabulum and the last specimen had irregularities in the piriformis fossa and femoral neck with abundant scar tissue.

6. DISCUSSION

Bifurcation of the femoral head is a rare entity, with few reports in the literature [1,2,4]. Our patient is the third to be reported who had prior surgery for DDH. In order to evaluate the hypothesis that surgical trauma during surgery for DDH may be related to the development of a bifid femoral head, an animal model was created. A bifid femoral head was demonstrated in 2 of 6 rabbits who underwent surgical bisection of the femoral head. Another specimen was similar on radiographic findings, however the histology revealed only one femoral head with a compact bone spicule similar to an exostosis.

The question remains: why did not all of the femoral heads become bifid post-surgery? The specimens without pathology in the femoral head may have been caused by surgical error. Additionally, the ossific nucleus may not have formed in 13 - 15 days since it has been shown by Heikel [6] that the rabbit ossific nucleus presents in only 43 of 57 animals at 6 - 10 days. Alternatively, the surgical insult may have healed and remodeled.

7. CONCLUSIONS

Bifid femoral heads can be induced by splitting the ossific nucleus in immature rabbits. This should be kept in mind when performing surgery around the ossific nucleus in immature patients. Focal necrosis and tight surrounding structures are not the etiology of the true bifid femoral head.

In children in whom a bifid femoral head is identified, a history of surgery at a young age, for DDH, should be sought.

REFERENCES


Figure 5. Gross specimen of double femoral heads and acetabuli.

Figure 6. Gross section: two femoral heads with articular cartilage, single physis.

Figure 7. Histology of two separate femoral heads, single physis.
necrosis of the femoral head as a complication of the dislocation of the hip in young children. A clinical and experimental investigation. Canadian Journal of Surgery, 12, 44.

