Rare Association of Bilateral Fracture of the Femoral Neck and Slipped Capital Femoral Epiphysis in Osteopetrosis Patient

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Abstract

We report a rare case of a five-year-old female patient who has Autosomal Dominant Osteopetrosis associated with bilateral fracture of the femoral neck (Salter & Harris II) and atypical mild slipped capital femoral epiphysis after running and playing, being initially treated conservatively with cast spica for 8 weeks, consolidating the right hip in varus, with decreased mobility ranges and positive Trendelenburg test. When the patient was 7-years-old, it was performed a right valgus femoral osteotomy with a plate screw system plus cast spica for 8 weeks, radiographs showed improved of neck shaft angle, no evidence of avascular necrosis in the affected femoral head, without delayed or non-union in the subsequent visit, integrating rehabilitation and improving ranges of passive and active mobility, as well as gluteal insufficiency. At five year follow up (12-years-old), the patient denies pain and presents good ranges of mobility.

Osteopetrosis is a rare bone dysplasia due to an alteration of osteoclast function in which bone resorption is decreased, resulting in alterations in bone mineral density, frequent fractures and risk of non-union or avascular necrosis, being an atypical case of bilateral femoral neck fracture plus slipped capital femoral epiphysis in a patient with Osteopetrosis. This rare association has never been reported.

Keywords: Children; Coxa vara; Femoral neck fracture; Osteopetrosis; Salter and Harris fracture; Slipped capital femoral epiphysis

Introduction

Osteopetrosis, also called marble bone disease, Albers-Schonberg disease belongs to the group of bone dysplasias, comprising a clinically and genetically heterogeneous group of conditions that share the hallmark of increased bone density on radiograph. The increase in bone density results from abnormalities in osteoclast differentiation or function and persistence of calcified chondroid material, causing the bone to be dense and brittle [1].

Albers-Schonberg was the first to describe in 1904 in a 26-year-old patient [1,2]. The decrease in osteoclast activity also affects the shape and structure of bone by altering its capacity to remodel during growth. In severely affected patients, the medullary cavity is filled with endochondral new bone, with little space remaining for hematopoietic cells. This abnormality contributes to the brittleness of bone in osteopetrosis.

This rare inherited disorder usually occurs in childhood or adolescence, although its diagnosis is also made at ages between 20 and 40 years.

Several forms of presentation of the disease have been described, but in all, the pathophysiological mechanism is the same, the lack of bone resorption due to a decrease in the number or activity of osteoclasts.

Osteopetrosis has been classified in three clinical forms [3], depending on the severity of its manifestations and the age of onset of the process:

- Childhood or malignant osteopetrosis. It is the most severe type and is inherited in autosomal recessive pattern. Its appearance is early in childhood and is characterized by a severe insufficiency of the bone marrow that usually has a poor prognosis.
- Benign or adult osteopetrosis. It occurs later, in young adults and is inherited with autosomal dominant character. In this variety they have a full life expectancy and there is no severe spinal insufficiency. There is a greater susceptibility to orthopedic pathologies such as fractures, osteomyelitis and early coxofemoral osteoarthritis due to fractures at the level of the femoral neck or slipped of the epiphysis, additionally sometimes some cranial nerves are affected by bone invasion in the holes at the base of the skull.
- Intermediate osteopetrosis. It is less serious than the infant variety and usually affects children under 10 years. It is transmitted recessively and is characterized clinically by cerebral calcifications, renal tubular acidosis and bones with increased density. In this rare variant, a deficiency of the carbonyl anhydrase II enzyme, necessary for osteoclast bone resorption, has been demonstrated.

One of the most serious complications of osteopetrosis is the femoral neck fracture, which has been rarely reported and very few studies about this case are present. Therefore, the optimal management of femoral neck fractures in patients with osteopetrosis has not been clearly defined. No validated treatment guidelines exist [4,5].
There is even less information available, when the injury is accompanied by bilateral fractures and Slipped Femoral Capital Epiphysis, related to potential complications including suboptimal healing, avascular necrosis of the femoral head, nonunion, post-traumatic coxa vara, growth arrest, infection, and chondrolysis [6,7].

We report a case of Osteopetrosis (ADO) with bilateral femoral neck fractures (Salter and Harris II) plus slipped capital femoral epiphysis treated in first round conservatively and then surgically as-yet-unreported.

**Presentation Case**

A 5-year-old female patient who came to the clinic of Hip Pathology with initial complaints of right painful weight bearing and limp for the last 2 weeks, and then bilateral pain as noticed by her mother after she was running and playing. As a significant antecedent, she got a fracture in the distal phalanx of the 5th right toe at one year of age and a right elbow fracture at 4 years old with conservative treatment. Her physical body appearance at 9 years and 5 months of age was normal with the only evidence of pectus carinatum. Height (51.18”) and weight (65.69 lb.) in average rank, Body Mass Index (BMI) was 17.6 (This indicates a BMI by age, that represent the 68th for normal children at the age of 9 years 5 months which means that she had a healthy weight at that period of evaluation. Dependent gait of crutches, with difficulty, painful to passive mobilization, a slight limitation of abduction and internal rotation. Laboratory blood tests showed 4.53 million cells erythrocyte count per microliter (mcL), Hemoglobin 12.4 g/dL, increased the levels of erythrocyte sedimentation rate in 35 mm/h, and slight increased white cell count (11.6 uL) predominantly neutrophils (62.3%) vs lymphocytes (28.7%).

Pelvis radiology and Panoramic x-ray of lower limbs were performed, showing extraordinary bone radiopacity, dense and faint medullary space, thickness of cortical and cancellous bone, bilateral large wedge-shaped fracture of the inferior metaphyseal border of the femoral neck (Red arrow in Figure 1), with inferior subluxation of capital epiphysis. The line of Klein (Red lines) describes a line along the superior edge of the neck of the femur. It is useful in detecting SCFE. A Southwick angle of 25º right and 24º based on femoral epiphyseal-diaphyseal angle difference in a frog-leg lateral x ray used to measure the severity of a slipped Epiphysis.

**Figure 1:** At 5-year-old. Note the extraordinary bone radiopacity, dense and faint medullary space, bilateral fracture of the inferior metaphyseal border of the femoral neck in Anteroposterior (AP) view (Red arrow), with inferior subluxation of capital epiphysis. The line of Klein (Red lines) describes a line along the superior edge of the neck of the femur. It is useful in detecting SCFE. A Southwick angle of 25º right and 24º based on femoral epiphyseal-diaphyseal angle difference in a frog-leg lateral x ray used to measure the severity of a slipped Epiphysis.

Treatment options (In situ pinning vs spica cast), prognosis, and complications were discussed with the parents. They were concerned about the surgery and therefore asked for the patient to be managed conservatively. The patient was initially treated with bilateral long leg hip spica cast (30º of hip flexion and abduction, 45º knee flexion) for 8 weeks. The other radiographs of spine, thorax and skull (Figure 2) showed increased sclerosis of the bone, vertebral end-plate thickening.

**Figure 2:** A lateral view of the thoracic spine showing increased sclerosis of the bone in the skull, ribs, and vertebral end-plate thickening.

X-ray at two months (Figure 3) show consolidating of the left fracture, and partial consolidation in the right side with coxa vara (100º right neck shaft angle vs 120º left hip) and stabilization of SCFE. Clinically no pain, right hip mobility ranges: Abduction (30º), Adduction (10º), Flexion (85º), Internal rotation (30º), External rotation (40º) and positive right Trendelemburg test. Limited weight-bearing was recommended for a lengthier period.

**Figure 3:** AP radiographs showing Consolidating of left Fractures, and partial consolidation in the right femoral neck with coxa vara (100º right neck shaft angle vs 120º left hip) and stabilization of SCFE.

When the patient was 7-years-old, a right valgus subtrochanteric osteotomy of the femur was performed, using a plate screw system for fixation. The patient had to wear a cast Spica for 8 weeks to help to consolidate, improving in the neck shaft angle of 122º (Figure 4).

**Figure 4:** Figure 2: A lateral view of the thoracic spine showing increased sclerosis of the bone in the skull, ribs, and vertebral end-plate thickening.

Subsequent radiographs showed no evidence of avascular necrosis in the affected femoral head, without delayed or nonunion, capacity of remodeling of spherical head and neck femur being congruent in the coxofemoral joint in successive consultations. There was no limp or pain, integrating rehabilitation and improving ranges of passive and active mobility (130º flexion, 45º abduction), as well as gluteal insufficiency.

At 9-years-old, she was scheduled to have removed osteosynthesis material. At the three years follow up (12-years-old) the child was...
not in pain or limp, and presented good ranges of mobility, only with shortening of 0.4 cm right femur (Figure 5 and Figure 6).

![Figure 4: Right valgus subtrochanteric osteotomy of the femur using a plate screw system at 7 years old in the AP and frog-leg lateral radiographs.](Image)

![Figure 5: At 12-years-old, AP and frog-leg lateral radiographs. Patient without osteosynthesis material, showed no evidence of avascular necrosis in the femoral head, without delayed or nonunion, and capacity of remodeling of spherical head femur being congruent in coxofemoral joint.](Image)

![Figure 6: Scanometry of lower extremities with shortening of 0.4 cm of right femur.](Image)

Discussion

The purpose of this paper is to show a different clinical presentation and the available options to face the natural complications of Osteopetrosis.

We did a review and investigated reports describing association of neck femoral fractures as consequence of Osteopetrosis (ADO) [5-8], or atypical Slipped Capital Femoral Epiphysis with ADO [9-11].

There was little the literature, and only one publication of the three pathological associations [12], no articles were found related Salter Harris II in proximal femoral fracture in Osteopetrosis, and none being associated with bilateral fracture of the femoral neck and atypical SCFE in ADO.

Fractures of the proximal third of the femur are rare in children and represent < 1% of all pediatric fractures. Most injuries result from high-energy trauma, such as a fall from a height or a car accident [4], unlike fractures in patients with endocrinological or metabolic conditions, such as in patients with osteopetrosis, where simple or minimal traumas of very low energy can cause fractures. Events without any traumatic history can trigger a stress fracture, especially in bones that support loads like the femur.

Krieg, et al. [6] support the surgical option as a recommended treatment especially in the neck of the femur, avoiding or decreasing the risks of non-union, coxa vara and subsequent avascular necrosis. The management of stress fractures of the femoral neck in pediatric patients with ADO remains controversial. Surgical and non-surgical treatments present their own challenges.

The Slipped Capital Femoral Epiphysis (SCFE) is a disorder of the child and adolescent hip that can be classified as idiopathic or atypical. Although the vast majority are idiopathic and are reflected in most scientific studies, only a small number of SCFE are atypical, and are those associated with endocrine pathologies such as thyroid disorders, renal failure osteodystrophy, metabolic disorders, osteopetrosis, or undergoing radiation, with little literature in the web, and increasing the risk of being unstable, with subsequent avascular necrosis, and worse prognosis [13,14].

Children presenting with SCFE who are younger than 10 years old or older than 16 years, whose weight is < 50 percentile for age, or whose height is < 10 percentile for age should be considered atypical. In this situation it should be mandatory to make a screening and investigate the atypical conditions mentioned above [15]. This is very important to consider as a differential diagnosis, there being no clear and defined guidelines about the management of this group of atypical disorders in the capital femoral epiphysis.

Anatomic factors such as obesity, proximal femoral, retroversion, and a vertical physis are known elements contributing to the mechanical pathogenesis of SCFE. The goals of SCFE surgery are to achieve stabilization, Avoid Avascular Necrosis (AVN), and minimize limitations in function [14].

Kinig, et al., [9] indicate four factors have been postulated in the pathogenesis of slipped capital femoral epiphysis: Increased height of the epiphysis, changes in the inclination angle of the physis, abnormal loading of the growth plate, and insufficiency of the components of the physis cartilage (collagen and proteoglycan). In the case of patient with Osteopetrosis there is a metabolic defect in the resorption of the bone being prone to multiple microfractures due to defective remodeling increasing the stress and shearing forces across the growth plate that could extend to metaphysis like in our patient. To Abaci, et al., [10] the changing the distribution of weight bearing on the joints is a factor in developing SCFE.

Dharmshaktu, et al., [12] shares a case of a 12-year-old male similar to ours, but in only one side with apparent inferior subluxation of...
capital epiphysis on right side along with cortical irregularity along neck region. The very presence of decreased neck shaft angle may also increase chances of neck fracture due to higher strain and shear force concentration.

The dense structure of osteopetrosis bone is associated with a low modulus of elasticity, allowing little bony deformation, increasing the risk of tension fractures, coxa vara and degenerative arthritis over time [16-18].

Armstrong, et al., [18] reported in a survey of 172 members of Pediatric Orthopaedic Society of North America (POSNA) the largest report of femoral neck fracture in patients with Osteopetrosis. 6 patients aged 6 years to 16 years with 7 cases of acute femoral neck fracture were reported. Four of those fractures were treated with closed reduction and internal fixation by using either pins or compression screws without complications. The others 3 were managed nonoperatively with posterior complications like non-union and varo, requiring valgus osteotomy. Our case was in a 5-year-old girl with a bilateral fracture of the inferior metaphyseal border of the femoral neck (Salter Harris II) who secondarily displaced both epiphyses, being the youngest patient with this type of rare complication with unilateral coxa vara that required subtrochanteric osteotomy.

Although coxa vara is a well-known complication of osteopetrosis, there are few reports of its treatment, being difficult to correct surgically due to the severe deformity because of the hardness and brittle character of the bone substance, increasing the risk of fail and complications. Most of these cases had gradually developed the deformity during early childhood and had become symptomatic with pain and or limping, being treated with a valgus Osteotomy [17,18].

A period of non-weight bearing, and bed rest is indicated because of the greater risk of delayed union or nonunion in these patients [18,19]. As is pointed out by Gunther, et al. [20] pathogenetically a coxa vara mostly emerges out of a biological insufficiency of the femoral neck or a disturbance of the epiphyseal growth plate, both of which are incapable of handling force loads of everyday activities.

The result of coxa vara and the consequent overgrowth of the greater trochanter bring the hip abductors’ sites of origin and insertion closer together, which results in decreasing tension and efficiency of glutaeus muscles to keep the pelvis stable, resulting in considerable lordeosis of the lumbar spine, clinically showing in a positive Trendelenburg’s sign with shortening of the affected leg [20], presenting in our patient.

Decreasing the angle of the neck axis can also increase the chance of neck fracture due to a higher concentration of tension and shear force in patients with osteopetrosis [6] and increase the risk of degenerative arthritis over time [16,17,19].

In a study by Palma, et al. [21], bone callus formation was histologically evaluated in a patient with autosomal dominant osteopetrosis (OAD) at different times, revealing normal bone tissue formation at 20 days of a fracture, with normal osteogenesis, with woven bone and occasional cartilage islands with fewer vessels with the characteristics elastics fibers and deformed with scant cytoplasmic osteoclasts. After one year of evolution, a consolidated fracture biopsy was performed, showing primitive bone tissue with lack of lamellar organization, with the absence of osteoclasts. Thus, osteopetrotic fractures do heal, but remodeling does not occur. Moreover, removal of the plate and screw can cause a new concentration of stress corresponding to the screw holes, which can be considered permanent sites of lower resistance, and risk of new fracture.

Little difficulty was experienced with perforation of dense bone in our patient and applying the osteosynthesis, probably because of the mild degree of the disease process. To decrease the risk of thermal bone necrosis [22-24], we always try to avoid the use of a power drill in performing a medullar cervical canal, using manual reamed over a guide wire, and power drill for cortical holes.

Our publication of the bilateral fracture neck femur and SCFE in ADO complicated with coxa vara, is the first report in documented literature, highlighting the importance of categorizing, reporting, and giving alternatives of treatment, the need for close follow-up the patient, and management of these cases with high risk of complications. The use of the screw plate system for the fixation of the proximal femoral valgus subtrochanteric osteotomy proved to be useful in the stability of the proximal femoral third by restoring the neck shaft angle and biomechanical of the hip.

Conflicts of Interest

No benefits in any form have been received or will be received from a commercial party related directly or indirectly to the subject of this article.

References


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