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Case Report

Resection and Reconstruction of Fibrocartilaginous Mesenchymoma of the Distal Fibula: A Case Report and Review of the Literature

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Abstract

Fibrocartilaginous mesenchymoma (FM) is a rare, locally aggressive bone tumor, with only 37 cases have been reported in the literature since 1984. This tumor principally occurs in long metaphysis bones in children and adolescents. In this case, a 13-years-old boy presented to the hospital with a firm, immobile, painful and slow growing mass of the right ankle that had been apparent for 4 months. The pathology after the biopsy showed Fibrocartilaginous mesenchymoma. We performed right fibula distal tumor resection and vascularized fibula reverse graft to repair the bone defect, which not only completely resected the tumor, but also preserved the stability of the ankle joint and ensured blood supply of the fibula reverse graft. The ankle joint function was stable in the short-term follow-up after operation.

Keywords: Fibrocartilaginous mesenchymoma; Tumor of the distal fibula; Bone tumor; Vascularized fibular graft; Resection; Reconstruction

Introduction

Fibrocartilaginous mesenchymoma (FM) is a rare, locally aggressive, primary intraosseous borderline tumor with unknown etiology, all of which are single, often asymptomatic or only manifested as pain or swelling at the lesion site [1]. Radiologically, FM appears as an expansible osteolytic lesion with cartilaginous calcification and cortical destruction, and extension to soft tissue is not uncommon. Histologically, FM is characterized by spindle cell proliferation in association with bland cartilage nodules and epiphyseal growth platelike enchondral ossification [2].

Fibrocartilaginous mesenchymoma was first reported by Dahlin et al. in 1984 [3]. A total of 37 cases of FM have been reported in foreign literature (Table 1). In all reports, patients range in age from three months to 27 years (median age of 13 years) at the onset of symptoms. Among them, there are 25 males and 12 females (male: Female=2.08:1.00), this tumor occurs in long metaphysis bones (21 cases), other locations are the iliac-pubic bone (6 cases), vertebrae (6 cases), ribs (2 cases), and metatarsal (1 case). All cases were followed up for 4 to 196 months, and 4 cases relapsed after partial resection. In this case, the patient was followed up for 9 months after the operation, and there was no sign of recurrence.

Case Presentation

A 13-years-old boy presented to Henan Provincial People's Hospital (Henan, China) with a firm, immobile, painful and slow growing mass of the right ankle that had been apparent for 4 months, the pain was aggravated during activity. A physical examination revealed there was a long oval mass on the right ankle, which was firm, immobile, painful, no local redness and local skin temperature was not high. Laboratory examination of alkaline phosphatase (ALP: 466.0 U/L) was higher than the normal value. X-ray (Figure 1A) revealed: a cystic low-density shadow could be seen at the distal end of the right fibula, with a clear boundary, no obvious sclerosis zone around it, and swelling growth. There was no obvious abnormality in the adjacent bone marrow cavity, and the local bone cortex was thinning and defect; multiple atrial septals were seen in the lesion, and no obvious periosteal reaction was seen. Computed tomography (CT) (Figure 1B) revealed that the distal end of the right fibula showed swelling changes, with patchy dense calcifications and cysts, and soft tissue density shadows, and the lesions did not cross the epiphyseal line. Magnetic resonance imaging (MRI) (Figure 1C) revealed that the swelling changes in the distal metaphysis of the right fibula showed long T1 and long T2 signal shadows, and the boundary was clear. Line-like long T2 signal shadows were visible in the surrounding soft tissues, and there was no more obviously abnormal. Emission CT (ECT) (Figure 1D) revealed that only the radioactive abnormal concentration was seen in the distal end of the right fibula, and there was no obvious abnormality in bone metabolism in the rest. The pathology of needle biopsy under local anesthesia showed that the possibility of fibrocartilaginous mesenchymoma was considered. It was recommended to confirm the diagnosis after tumor resection.

Operative steps

The tumor was approached through a direct posterolateral incision over the distal half of fibula. The cutaneous branch of superficial peroneal nerve was identified and preserved. The distal end of the tumor did not invade the epiphysis, the proximal end of the tumor was osteotomized along the epiphyseal line, the epiphysis was preserved (about 1cm), the proximal end of the tumor 3 cm outside was osteotomized, and the tumor was completely resected (Figure 2A). In order to ensure the blood supply of the fibula reverse graft, we found a nourishing artery (distal branch of the peroneal artery) at the normal fibula near the fractured end of the osteotomy, and measured the distance about 5cm from the distal branch of the peroneal artery

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Patient and Reference	Age (years), Sex	Location	Treatment	Further treatment	Follow up
1 [4,5]	10y F	Third rib	Wide resection	—	NED at 4y
2 [4,5]	9y M	First metatarsal	Intralesional excision	Wide contaminated resection of recurrence (1y later); then wide resection of a second recurrence (1y later)	NED at 12y
3 [4,5]	14y M	Metaphysis proximal fibula	Intralesional excision	Wide resection of recurrence (6y later)	NED at 78mo
4 [4,5]	9y M	Metaphysis proximal fibula	Wide resection	_	NED at 5y
5 [4,5]	22y M	L4	Intralesional excision	—	NED at 10y
6 [4,5]	14y M	Diaphysis fibula	Intralesional biopsy	Intralesional biopsy (1y later)	UK
7 [4,5]	14y M	Proximal tibia	Resection	—	NED at 5y
8 [4,5]	15y F	Pubis	UK	—	UK
9 [4,5]	25y F	Proximal humerus	Resection		NED at 2y
10 [4,5]	16y M	Proximal tibia	Intralesional excision	Resection of recurrence (2y later)	NED at 2y
11 [4,5]	12y F	Proximal Fibula	Resection	_	NED at 1y
12 [5,16]	11y M	Iliac bone	Incomplete excision, then wide resection	—	NED at 14y
13 [5,17]	4y M	Metaphysis proximal humerus	Curettage with adjuvant penalization	_	NED at 2y
14 [5,18]	17y F	Pubis	Intralesional excision with phenol and ethanol cauterization	—	NED at 1y
15 [5,19]	3mo F	Metaphysis proximal tibia	Biopsy	_	complete regression afte 14mo
16 [5,20]	11y M	Metaphysis proximal humerus	Forequarter amputation	_	NED 2y
17 [5,21]	9y M	T12	Resection	—	NED 2y
18 [5,22]	15y F	Metaphysis distal femur	Resection and curettage of a part of the tumor	_	NED 5y
19 [5,23]	9y M	Metaphysis distal femur	Wide resection	_	NED 4y
20 [5,24]	9y M	llium	UK	—	UK
21 [5,25]	14y M	Metaphysis proximal humerus	Resection		UK
22 [5,26]	19y M	L5	Wide contaminated resection	_	NED 5y
23 [5,27]	12y M	Metaphysis proximal tibia	Curettage	Wide resection of recurrence (1y later)	NED 2y
24 [5,27]	1y 7mo M	Metaphysis proximal tibia	Curettage		Decreased in si after 4mo
25 [5,28]	11y M	Metaphysis proximal tibia	Curettage, then wide resection	_	NED 10y
26 [5]	8y F	Metaphysis proximal femur	UK	_	UK
27 [5]	10y M	Distal femur	Intralesional excision	UK treatment of recurrence (1y later)	UK
28 [5]	27y F	Pubis	Resection	_	NED 17mo
29 [5]	18y F	llium-pubis	Resection and curettage of a part of the tumor	_	NED 65mo
30 [5]	18y F	L3	En bloc wide resection	—	NED 10y
31 [5]	13y M	L4	En bloc wide resection	_	NED 38mo
32 [5]	14y M	Metaphysis proximal tibia	Intercalary resection	_	UK
33 [5]	22y F	Metaphysis proximal humerus	Curettage	wide Tikhoff-Linberg resection (2 weeks later)	NED 196mo
34 [1]	13y M	L3	Wide resection	—	NED 9y
35 [29]	16y M	Metaphysis proximal tibia	En bloc resection	_	NED 7mo
36 [2]	17y M	Fifth rib	En bloc resection		NED 1y
37cs	13y M	Distal fibula	En bloc resection		NED 6mo

Table 1: Location, clinical features, further treatment, and follow-up.

Legend: y: year(s); mo: months; UK: Unknown; M: Male; F: Female; NED: Not Evidence of Disease; cs: current study.

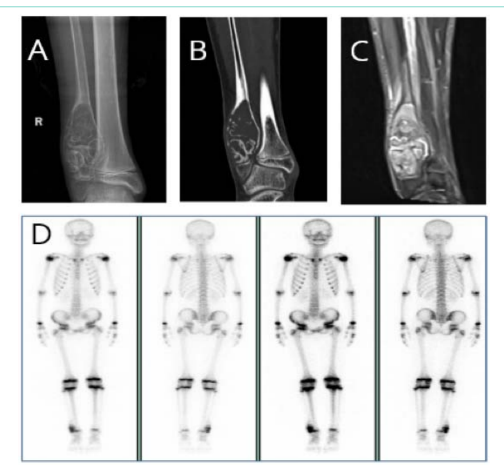


Figure 1: A) X-ray revealed that a cystic low-density shadow could be seen at the distal end of the right fibula, with a clear boundary. B) CT revealed that the distal end of the right fibula showed swelling changes, with patchy dense calcifications and cysts, and soft tissue density shadows, and the lesions did not cross the epiphyseal line. C) MRI revealed that the swelling changes in the distal metaphysis of the right fibula showed long T1 and long T2 signal shadows, and the boundary was clear. Line-like long T2 signal shadows were visible in the surrounding soft tissues, and there was no more Obviously abnormal. D) ECT revealed that only the radioactive abnormal concentration was seen in the distal end of the right fibula, and there was no obvious abnormality in bone metabolism in the rest.

to the distal epiphyseal line of the fibula epiphysis, and osteotomized the normal fibula from the distal branch of the peroneal artery to the proximal end of the normal fibula at about 5cm and dissected the fibula periosteum. The vascularized fibular graft was inverted, and we reconstructed the proximal fibula osteotomy surface and the distal epiphysis osteotomy surface (Figure 2B). We reconstructed the inverted vascularized fibular graft with a titanium plate and screws. X-rays showed that the internal fixation was firm and ankle joint was stabilized (Figure 2C).

The ankle was immobilized in a below knee back slab till suture removal. At suture removal, an ankle brace was given and patient was allowed intermittent ankle ROM as tolerated and touches weight bearing with bilateral axillary crutches for 6 weeks progressing to full weight bearing on ankle brace for another 6 weeks. Till 4 months, he was on protected weight bearing with ankle brace. Clinical and radiological follow up was done regularly. At 9 months follow up, clinical assessment was done with American Orthopaedic Foot and Ankle Society score and radiological assessment. At follow up at 9 months, there was no ankle instability. The AOFAS score was excellent (92/100). His dorsi flexion was 20° and plantar flexion was 40°.

The pathology (Figure 2E) is generally gray and white matter or fibrous tissue with tough texture, and there are scattered and distributed translucent and brittle cartilage nodules. The photomicrograph image (Figure 2D) shows that a large number of spindle cell proliferations are seen between the well-differentiated cartilage islands, the tumor cells are slightly atypia, mitotic images are occasionally (0-1/10HPF), and no tumor necrosis is seen. Localized cartilage matrix calcification and ossification, scattered multinucleated giant cells, and part of bone trabecula were seen around. Combined with immunohistochemical markers, it was consistent with fibrocartilage mesenchymal tumor; immunohistochemical results showed: -A1:B-Catenin (-), Bcl-2 (scattered +), Bcl-6 (-), CD31 (-), CD34 (-), CDK4 (-), CK (AE1/AE3) (-), CK19 (-), CK7 (-), Desmin (-), Ki67 (5%+), MDM2 (+), MSA (-), MyoD1 (-), Myogenin (-), Napsin A (-), Nestin (-), P16 (+), P53 (-), S-100 (cartilage island +), SMA (-), STAT6 (-), Villin (-), Factor VIII (vessel +), H3F3AG34W (-), SATB2 (+). FISH result: It is indicated that the MDM2 gene of the sample submitted for inspection was not amplified (negative). IDH: The IDH1/IDH2 gene mutation test result of the sample submitted for inspection is no mutation (wild type).

Discussion

Fibrocartilage mesenchymal tumor is a rare primary intraosseous

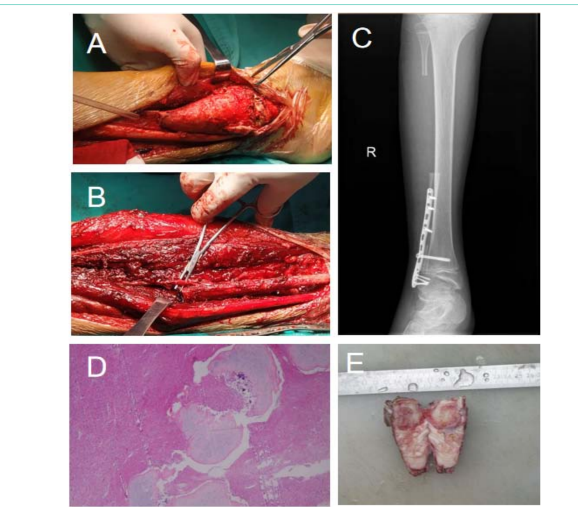


Figure 2: A) Exposing and removing the tumor of distal fibula. B) Inverting the vascularized fibular graft and reconstructing the proximal fibula osteotomy surface and the distal epiphysis osteotomy surface. C) Reconstructing the inverted vascularized fibular graft with a titanium plate and screws. X-ray examination showing that the internal fixation was firm and the ankle joint was stable. D) Pathological photomicrographs showed a large number of spindle cell proliferations between well-differentiated cartilage islands, mildly atypia of tumor cells, occasional mitotic images, no tumor necrosis, and focal cartilage matrix calcification and ossification. Scattered multinucleated giant cells can be seen, and some trabecular bones are seen around. E) The pathology is generally gray and white matter or fibrous tissue, with translucent and brittle cartilage nodules scattered in the middle.

borderline tumor [1]. In 1984, Dahlin et al. reported 5 cases of FM for the first time. Among them, 3 cases of local recurrence were named "low-grade malignant tumors" [3]. In 1993, Bulychova et al. retrospectively analyzed the case data of 12 FM patients and believed that incomplete resection could lead to local tumor recurrence, but no metastasis or death, so it could not be classified as "low-grade malignancy". "Follow-up case analysis and follow-up showed that there were few recurrences and metastases after complete resection of the lesions, which further proved this view [4]. The 5th edition of the World Health Organization classification of bone tumors newly listed FM as an independent tumor entity.

Radiologically, FM appears as an expansile osteolytic lesion with cartilaginous calcification and cortical destruction, and extension to soft tissue is not uncommon. Histologically, FM is characterized by spindle cell proliferation in association with bland cartilage nodules and epiphyseal growth plate-like enchondral ossification. The differential diagnoses include FCD (fibrocartilaginous dysplasia, FCD), low-grade osteosarcoma, dedifferentiated chondrosarcoma, desmoplastic fibroma, and chondromesenchymal hamartoma of the chest wall [2]. In 2017, Gambarotti M, et al. [5] analyzed eight new cases from the files of the Istituto Ortopedico Rizzoli dating from 1982 to 2016. This very rare bone tumor has a typical radiological and histological pattern and a favorable survival outcome after treatment. Local recurrences can be prevented with complete surgery. FM does not seem to be genetically related to fibrous dysplasia, low-grade osteosarcoma, and dedifferentiated chondrosarcoma.

Tumors of the distal fibula are rare. However, their management poses significant challenges. Given the low incidence and diverse nature of distal fibular tumors, there are several solutions [6]. For instance, distal fibular resection without reconstruction of the lateral side of the ankle is frequently performed [7]. In such instances, ankle stability is obtained via either soft tissue and ligament reconstruction or tibiotalar arthrodesis. In other cases, fibular resection is followed by reconstruction with allograft, autografts, pedicled vascularized Gao S

epiphyseal transfers using the ipsilateral proximal fibula or a long bone graft from the iliac crest, bone transplants, or prosthetic ankle joint replacement [8-10].

Different techniques for reconstruction of the distal fibula after wide tumor resection have been described. All methods have different advantages and disadvantages [11]. In this case, the tumor did not attack the distal fibula epiphysis and epiphysis line, and the child's epiphysis was not closed. If the distal fibula bone was completely resected directly, the loss of distal epiphysis would not only lose the opportunity of joint development, but also cause the loss of ankle joint stability. Resection of the lateral ankle can cause varus instability or a collapse into valgus [12]. If the distal fibula is resected and reconstructed with ipsilateral fibulae capitulum, loss of the proximal fibula can cause lateral knee instability or a damage of the peroneal nerve [13-15]. Another disadvantage of this technique is the incongruity of the fibula head with the articulating talus and the risk of pseudarthrosis. In addition, the distal branch of the peroneal artery which could be used as an effective blood supply source for the fibular graft ensured the blood supply of the inverted fibular graft and avoided the undesirable healing of the broken end of the fibular graft and the distal epiphysis of the fibular graft due to poor blood supply.

In summary, bone FM is a rare intraosseous borderline primary tumor characterized by fibrous hyperplasia and epiphyseal cartilage formation. FM is a locally aggressive tumor with no distant metastasis reported thus far. Local recurrence occurs only in cases of incomplete removal, such as curettage or intralesional excision. Complete surgical excision with adequate margins is the treatment of choice for FM.

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