Chondrosarcoma of the Foot: A Rare Occurrence in the Distal Phalanx of the Fourth Digit

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Abstract

Chondrosarcomas are among the most common types of tumors seen in the body. However, these lesions are very rarely found in the foot as there are few documented cases reported in literature. In this article, we present a case study of a primary chondrosarcoma arising in the foot involving the distal phalanx of the right 4th toe in a 38-year-old female. Initial radiographs did not reveal a fracture; however, slight abnormalities were seen in the bone that led to a high suspicion of a bone tumor. MRI confirmed the presence of neoplasm. Subsequently, a bone biopsy was performed with an intra-operative pathology consult. Pathology results revealed and intermediate-grade chondrosarcoma of the distal phalanx of the right fourth toe. The patient later went on to have an amputation of the right 4th toe at the level of MTPJ. Upon follow-up, a CT scan of the chest, abdomen, and pelvis was ordered to ensure that the spread of metastatic disease was prevented. The patient went on without progression of disease after 4 years.

Introduction

Chondrosarcomas are the third most common malignant tumors of bone, being less common than only myelomas and osteosarcomas. Chondrosarcomas are a group of tumors with highly diverse features and behavior patterns, ranging from slow-growing non-metastatic lesions to highly aggressive metastasizing sarcomas [1]. Chondrosarcomas are malignant tumors of cartilaginous origin in which tumor matrix formation is entirely chondroid in nature. Chondrosarcomas can be divided into several different subtypes depending on clinical, radiographic, and histologic characteristics. The subtypes are: conventional chondrosarcoma, chondrosarcoma of small bones, secondary chondrosarcoma, perioskeletal chondrosarcoma, clear cell chondrosarcoma, dedifferentiated chondrosarcoma, and mesenchymal chondrosarcoma [2].

Chondrosarcomas are most frequently found in men between the ages of 30-60 years although there is a slight female predominance for phalangeal chondrosarcomas. Chondrosarcomas are typically slowly evolving neoplasms. The most common symptoms are mild pain and tenderness. Other physical findings include localized swelling, well defined soft tissue masses, pathologic fractures, and/or antalgic gait. The risk factors for chondrosarcoma include enchondromas, multiple exostoses, Ollier’s disease and Maffucci’s syndrome. Metastasis of chondrosarcoma from the foot is through a hematogenous route and is typically to the lungs. The overall prognosis is related to the size of the lesion, its anatomic location, and its histologic grade. Patients with axial lesions have a worse prognosis than those with lesions of the appendicular skeleton. The 5 year survival rate for patients with grade 1 lesions is 90%; the rate decreases to 29% with grade 3 tumors. Grade 1 lesions do not metastasize. Metastatic spread, typically pulmonary, is more frequently associated with grade 3 lesions than with other grades.

The radiographic appearance of these tumors varies, but conventional chondrosarcomas are usually seen on radiographic examination as large radiolucent areas with thick walls. These radiolucent areas have trabeculation with multilocular medullary bone destruction in the central areas. Secondary chondrosarcomas may have features that indicate the transformation of a benign osteochondroma into a malignant chondrosarcoma. These features usually consist of a bulky cartilaginous cap (more than 2 cm thick), scattered calcifications in the cartilaginous part of the tumor, focal areas of radiolucency, significant soft-tissue mass, pressure erosions or adjacent bone, and rapid growth. The most significant radiographic findings that indicate whether a tumor is a low grade or high-grade tumor are the pattern of calcification, the nature of the tumor margin, and the size of the soft-tissue mass. The pattern of calcification of low-grade tumors is widespread but without soft-tissue extension. There is a small zone of transition between normal and abnormal...
appearing tissue. High-grade tumors also have large areas with no calcification, as well as soft-tissue extension.

Low-grade chondrosarcomas may not be visible on radiographs, or they may show few or no signs of malignancy. It is important, therefore, to consider the use of other imaging modalities such as magnetic resonance imaging (MRI) or computed tomography (CT). MRI allows complete visualization of the intra-osseous and soft-tissue extent of the chondrosarcomas. Cartilaginous areas of the tumor have intermediate signal intensity relative to adjacent skeletal muscle on T1-weighted images and high signal intensity on T2-weighted images. Calcifications appear as areas devoid of signal [3-6].

Computed tomography and standard radiographs are the best modalities for visualization of calcification. If only minimal calcification is seen on standard radiographs or CT scans, it will not be seen on MRI. MRI not only is the best modality for visualization of the soft-tissue extent of the chondrosarcoma, but is also best for observing endosteal scalloping and cortical erosion. MRI can be used to characterize tumor grade. When MRI is enhanced with gadolinium, it may be helpful in determining biopsy location by delineating areas of necrosis within the lesion.

These physical and radiographic findings help in distinguishing malignant and non-malignant tumors of the foot. The differential diagnosis of chondrosarcoma in the foot includes benign lytic bone tumors such as enchondroma or osteochondroma, malignant bone tumors such as those that have metastasized from other areas or osteosarcoma, and infection, such as mycetoma or tuberculosis. Conventional chondrosarcomas are divided into four histological grades based upon their appearance under a microscope. The grading is based primarily on nuclear size of tumor cells, nuclear staining, and cellularity. Grade I (low grade) tumors resemble normal cartilage, but may surround areas of lamellar bone, or show atypical cells including bi-nucleate forms. Grade II (intermediate grade) is more cellular with a greater degree of nuclear atypia, hyperchromasia and nuclear size. Grade III (high grade) tumors have significant areas of marked pleomorphism, large cells with more hyper chromatic nuclei than grade II, occasional giant cells and abundant necrosis. Mitoses are frequently detected. Mesenchymal and dedifferentiated Chondrosarcomas are considered to be Grade IV tumors. Dedifferentiated chondrosarcomas, along with mesenchymal Chondrosarcomas, are highly malignant, particularly aggressive (i.e. rapidly growing and disturbing surrounding tissues) and carry a poor prognosis.

Secondary chondrosarcomas arise in the presence of a pre-existing condition. These may arise in exostoses, either single or multiple, or in chondrodysplasias. Most Chondrosarcomas arising in an osteochondroma are extremely well differentiated and, hence, are difficult to diagnose histologically. The radiographic features are important. Osteochondromas have a thin, regular cartilage cap. When a chondrosarcoma supervenes, the cartilage cap becomes thicker and irregular with fuzzy borders. Most osteochondromas have a smooth cartilage cap that is usually less than 1 cm thick. A thick cartilage cap, especially one showing myxoid change, suggests chondrosarcoma. Most secondary chondrosarcomas are low grade. The prognosis in secondary chondrosarcoma is generally good. Periosteal chondrosarcomas occur on the surface of a bone. These tend to be large lesions, usually more than 5 cm in greatest dimension. The radiographs show a poorly defined mass with uneven calcification. Histologically, the tumor tends to permeate surrounding soft tissues. Clear cell chondrosarcoma is an unusual chondroid neoplasm. The lesion tends to occur at the ends of long bones, similar to chondroblastomas and giant cell tumors. The radiographic appearance may mimic that of a
chondroblastoma in that the lesion is usually well circumscribed and may even have a sclerotic border. Aneurysmal bone cyst-like changes often are found in clear cell chondrosarcoma. The clinical behavior of clear cell chondrosarcoma is that of a low-grade chondrosarcoma. Dedifferentiated chondrosarcoma occurs in older adults. The imaging studies show classic features of chondrosarcoma but, juxtaposed to it, there is a more destructive-appearing area. Dedifferentiated chondrosarcomas are of a low-grade chondrosarcoma. Juxtaposed to it is a soft, fleshy sarcoma-like tumor. Microscopically, one sees a low-grade chondrosarcoma juxtaposed to a high-grade spindle cell sarcoma. The spindle cell malignancy is always high grade and may have features of fibrosarcoma, osteosarcoma, or malignant fibrous histiocytoma. A poor prognosis is associated with a dedifferentiated chondrosarcoma.

Dedifferentiated chondrosarcoma has to be differentiated from chondroblastic osteosarcoma. Chondroblastic osteosarcoma usually involves adolescents, whereas dedifferentiated chondrosarcoma involves older adults. In chondroblastic osteosarcoma, the cartilage cells look malignant and merge into a spindle cell sarcoma. In dedifferentiated chondrosarcoma, the cartilage is well juxtaposed to it rather than merging into it. This distinction is important because the prognosis in dedifferentiated chondrosarcoma is much worse than in chondroblastic osteosarcoma.

Mesenchymal chondrosarcoma affects mainly adolescents and young adults. About one-third of mesenchymal chondrosarcomas occur in soft tissues or the meninges. Mesenchymal chondrosarcoma tends to involve the jaw bones and the ribs. The radiographic features are non-specific. The radiographs usually suggest a malignant tumor, with or without mineral. Grossly, the lesion usually is pink and fleshy but may show foci of calcification. The microscopic appearance of mesenchymal chondrosarcoma is typical. There is a combination of well-differentiated cartilage and small cell malignancy. The long-term prognosis of mesenchymal chondrosarcoma is poor. This article reports a case of an intermediate grade primary chondrosarcoma in the proximal phalanx of the right fourth toe.

Case Presentation

A 38-year-old woman presented to the office for evaluation and treatment of right foot pain for two weeks duration. She was doing an activity where she was standing on her “tip toes” and she felt a pop and had immediate pain and discomfort localized to the 4th toe. She also complained of sharp, shooting pain with every step. Her past medical history was unremarkable. Pain, swelling, ecchymosis at the MTPJ of the fourth toe, and an antalgic gait were noted on physical exam. X-rays did not demonstrate any fracture, but did show an expanding osteolytic lesion with calcification in the distal phalanx of the right fourth toe which was highly suspicious of a bone tumor. Initial treatment consisted of a walking boot and an MRI to assess the bone tumor (Figure 1).

Patient returned to office after getting an MRI. Findings on the MRI showed a lesion in the fourth toe proximal phalanx with low T1 and bright T2 signal. The lesion approximated 12 mm in length (Figure 2). The medial and lateral cortices were grossly intact. Subsequently, a bone biopsy was performed with an intra-operative pathology consult approximately one week after receiving the MRI results. Histological sections showed lobular lesions composed of a combination of myxoid matrix and chondroid.

The cells were spindle or stellate in shape, some bi-nucleated cells were also seen (Figure 3). The cellularity was much higher than in normal cartilage. Chondroid cells were atypical. Final report diagnosed the lesion as an intermediate grade chondrosarcoma. The patient was referred to oncology once pathological results returned. The patient was worked up clinically by oncology and noted not to have any masses or ipsilateral inguinal masses. Blood work consisting of BMP 21, complete blood count with differential, liver function tests and thyroid panel all returned within normal limits (Figure 4).

The patient underwent diagnostic testing inclusive of CT scans of thoracic, abdominal and pelvic areas post operatively, 6 months and 1 year and 5 years after diagnosis which did not reveal metastasis (Figure 5).

Two weeks after the biopsy, an amputation was performed at the level of the 4th MPJ. The patient was followed up post operatively weekly until the surgical site healed completely in 4 weeks. Sutures were removed 2 weeks post operatively (Figure 6). The patient then followed up with the surgeon every 3 months until one year post op and then went to one year follow up appointments. The patient was monitored every 6 months with oncology until one year post operatively and then yearly (Figure 7). Five years post operatively the patient remains to do well with no complications from surgery or recurrence of cancer (Figure 8).

Discussion

Chondrosarcomas account for 10% to 15% of primary bone tumors. The most common sites for these tumors are the pelvis, shoulder, and the metaphysis of long tubular bones, particularly the femur. The incidence of primary bone tumor in the foot is about 2%, chondrosarcoma is the most common primary malignant bone tumor of the foot. According to Harkless, 36% of the tumors occurred in the
calcaneus, 41% in the metatarsals, and only 19% in the phalanges [1-3]. Correct classification of the grade of chondrosarcoma is important because the prognosis and treatment are different for different stages and grades. Evans used their own grading system: low grade is a low-level malignancy, intermediate grade-as in our case-is moderately malignant, and high grade is highly malignant. Reported 10-year survival rates for chondrosarcoma were 83% for low grade, 64% for intermediate grade, and 29% for high grade. Local recurrence was 40% for low grade, 60% for intermediate grade, and 47% for high grade. The incidence of metastatic disease is rare for low grade, 10% for intermediate grade, and 71% for high grade [4-7].

Bovee reported that phalangeal chondrosarcoma recurs in 36% of cases but does not metastasize [8-10]. Phalangeal chondrosarcoma also has a much better prognosis than chondrosarcoma in other locations. Mohammadianpanah described a case of a phalangeal chondrosarcoma at the proximal phalanx of the third toe that went on to an amputation at the MPJ [11,12]. Lo reported on a case of a low-grade chondrosarcoma at the base of the fifth metatarsal, lateral aspect of the cuboid and third cuneiform. The patient went on to have a below knee amputation.

Conclusion

In this case, early diagnosis of chondrosarcoma before metastasis and treatment with definitive amputation led to a malignancy free patient at 5 years post operatively. The patient continues to do well and functions without difficulty. Proper collaboration between the surgeon, pathologist, oncologist, radiologists and other medical specialists led to, at this point and hopefully for her foreseeable future, a healthy patient.

References