Chondrosarcoma of Second Toe Distal Phalanx-A Case Report

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Abstract

Chondrosarcoma is a relatively common primary malignant bone lesion. It is a malignant, relatively slow growing cartilage producing tumor. It forms approximately 10% of malignant primary bone tumors. However, it usually arises in long bones and truncal bones, whereas these tumors are uncommon in the hands and feet. The type are (A) Primary chondrosarcomas arise denovo from bone without any preexisting lesion. (B) Secondary chondrosarcomas when tumor arises from preexisting benign lesions of bone. Primary tumor is generally seen between 40 to 60 years of age. Secondary chondrosarcomas are more common in young adults.

Keywords: Chondrosarcoma; Tumor; Chondroblastoma; Bone; Neoplasm

Pathology of Chondrosarcomas

Central type

It is a lobulated, translucent, bluish whitish cartilaginous mass with in medullary cavity. Endosteal aspect of cortex appears eroded and scalloped. Tumor spreads widely through medullary cavity, its extent is greater than it appears on X-rays. As tumor penetrates the cortex it is temporarily delimited by periosteum. New reactive new bone is formed at the periphery of sub periostealmass there by thickening the cortex. This is seen in slow growing and less malignant tumors.

In rapidly growing tumors periosteum is penetrated easily, so no reactive new bone formation. When a firm greyish tanned tissue without lobular configuration is found, it represents most malignant type. On removing the closely investing periosteum, the tumor appears lobulated, bluish white, shiny, opalescent or semitranslucent mass of firm texture or rubbery consistency.

The cut surface exhibits specks of calcification that impart a gritty sensation on palpation, gelatinous or myxoid areas and ragged cavities. The degree of calcification is greater in slower growing tumors.

Microscopic appearance

Histological features of cartilage should be noted to diagnose presence of a cartilaginous tumor. A normal chondrocyte is ellipsoid, ovoid or round in shape, has slightly scalloped surfaces. It contain a central, oval, sometimes indented nucleus with regularly distributed abundant chromatin and indistinct nucleolus.

Sarcomatous chondrocytes and their nuclei vary greatly in size, shape, number of nuclei and nucleoli. Histology shows signs of malignancy including hypercellularity, plumpness of nuclei, double nuclei in more than occasional cells, pleomorphism of cells and nuclei, hyperchromatism and mitotic figures. Mitotic figures are rare in low and medium grade tumors but are readily found in high grade tumors.

Radiological Features

Central tumors

The lesion is seen in metaphysic or diaphysis of a long tubular bone. Rarely it may occupy epiphysis at juxtaarticular area, where it may be mistaken for a giant cell tumor or chondroblastoma. The lesion is located in medullary cavity, it is large and ill defined. Countour of bone is enlarged or expanded. The cortex is focally or extensively eroded and thinned. The inner cortex may have scalloped borders. Tumors will have specks of calcification and cotton whorl appearance. Slow growing tumors will show a greater of calcification where as highly malignant rapidly growing tumors will show little or no calcification and often destroys the cortex without expanding shaft. There is a associated soft tissue mass.

Low grade features

Dense calcification forming rings or specules with eccentric lobular growth of a soft tissue mass

High grade features

Faint amorphous calcifications with large non calcified areas and concentric growth of soft tissue mass.

This article presents the case of a 60-year-old woman with a chondrosarcoma in the distal phalanx of the left second toe. Radiography showed bony mass extending from the distal phalanx. Histologically, the tumor was classified as grade 1 chondrosarcoma [1-6].

Case Report

• A 60 yrs old lady presented with pain and swelling of the left second toe for the past 1 year.
• Insidious onset and Gradually progressive
• No rest pain/night cries/constitutional features
• No other swellings in body

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• k/c/o Type 2 diabetics and systemic hypertension for the past 2 yrs on regular medication

Local examination (left 2nd toe)
• Irregular well defined Swelling over the distal phalanx 2 X 3cms.
• Minimal deep tenderness, No warmth.
• Heterogenous consistency ranging from firm to bony hard.
• Adherent to underlying soft tissues and Immobile.
• Non pulsatile, Non fluctuant and Opaque.
• ROM–full range (Figure 1).

Plan
Patient was taken up for excision biopsy under local anesthesia and using Ray's amputation technique disarticulation of the second toe at MTP joint (Figure 3).

Histologically
Sections show fragments of bone with an infiltrating neoplasm composed of nodules of cartilaginous tissue with varying cellularity (hypercellular). The chondrocytes have moderate amount of cytoplasm mildly atypical hyperchromatic nucleus and few showing prominent nucleoli. Binucleate and trinucleate chondrocytes are seen. The cartilaginous matrix appears thin and mucoid at places. The tumour seen permitting the cortical bone causing bone entrapment (Figure 4).

Discussion
Chondrosarcoma of the foot is rare. Correct classification of the grade of chondrosarcoma is extremely important because the prognosis and treatment are different for different stages and grades (Table 1). Evans et al. used their own grading system: low grade is a low-levelmalignancy, intermediate grade is moderately malignant, and high grade is highly malignant. The chondrosarcoma of the phalanx is rare but, when present, extirpation is necessary to prevent growth and spread. Therefore, every effort should be made to differentiate between enchondroma and chondrosarcoma in each case. Complete wide excision of the lesion and cartilage cap should not be violated during resection of chondrosarcoma arising out of osteochondroma as it will increase the risk of local recurrence.
**Prognosis:** Tumor arising in osteochondromas have excellent prognosis. Secondary chondrosarcomas arising in enchondromatosis have same prognosis as that of conventional chondrosarcomas [1-6].

| Grade                  | Tumor                        | Symptoms          | Prognosis | Treatment                                                                 |
|------------------------|------------------------------|-------------------|-----------|---------------------------------------------------------------------------|
| Benign                 | Enchondroma                  | Usually no symptoms | Excellent | Surveillance, intralesional excision if symptomatic                        |
| Benign                 | Osteochondroma               |                   |           |                                                                           |
| Malignant (Low grade)  | Grade 1 Chondrosarcoma       | 60% are painful   | Excellent | Controversial: Extended intralesional excision vs. wide resection         |
| Malignant (Low grade)  | Grade 2 Chondrosarcoma       | Up to 80% are painful | Good     | Wide resection                                                           |
| Malignant (Intermediate grade) | Grade 3 Chondrosarcoma | Up to 80% are painful | Fair     | Wide resection. Chemoetherapy and radiation therapy in selected cases     |
| Malignant (High grade) | De-differentiated Chondrosarcoma | Most are painful | Poor      | Wide resection. Chemoetherapy when possible in all cases. Radiation therapy in selected cases |
| Malignant (High grade) | Mesenchymal Chondrosarcoma   | Pain and swelling  | Fair-Poor | Wide resection. Chemoetherapy in all cases.                               |

**Table 1:** Classification [6].

**Conclusion**

Phalangeal chondrosarcoma behaves as locally aggressive lesion and rarely metastasize. Curettage with local adjuvant and adequate follow-up can be tried as the first-line management, especially where amputation would lead to significant loss of function.

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