A Narrative Review on the Characteristics and Treatment of Benign and Malignant Bone Tumors

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SUMMARY

Bone tumors, including benign and malignant lesions, are not metastatic; however, they may appear in any part of the body skeleton. Distal femur and proximal tibia (around the knee joint) are the most prevalent sites. Most benign bone tumors are cartilaginous tumors, known as osteochondromas. Based on the reports, benign bone tumors are more frequent than primary malignant ones. Malignant bone tumor is another type of bone tumor, which usually occurs within the first years of life. As a result, it can considerably affect the lives of patients and their families. These tumors consist of osteosarcoma, chondrosarcoma, and Ewing’s sarcoma. This article discusses the epidemiology, characteristics, and treatment of the most important types of benign and malignant bone tumors. These data will be useful to the physicians and other health workers to better understand the conditions of bone tumors and their management.

Key words: bone tumor, malignant tumor, benign tumor
EPIDEMIOLOGY AND CLASSIFICATION OF BONE TUMORS

Bone tumors are divided into two categories, which include primary (occurs in bone or bone-derived tissues) and secondary (occurs in other areas and may metastasize to the skeleton) bone tumors (1). Prostate, breast, lung, thyroid, and kidney carcinomas are typically metastatic to bone (2). The estimated prevalence of secondary malignant bone tumors is as high as 50 - 100 times the primary bone cancers. Primary bone tumors are divided into benign and malignant ones (3). Some types of bone tumors are benign, and may be progressive, cancerous, infectious, and inflammatory (4). Some other kinds of bone tumors are benign and not metastatic. Tumors may appear in any part of the body skeleton, but the most prevalent sites are the distal femur and proximal tibia (around the knee joint). They become hazardous if they grow in healthy bones. Based on the matrix produced by cancerous cells, these tumors are classified into eight types: osteochondroma, osteoma, osteoid osteoma, osteoblastoma, giant cell tumor, aneurysmal bone cyst, fibrous dysplasia, and enchondroma. Malignant bone tumors consist of osteosarcoma, chondrosarcoma, and Ewing’s sarcoma (5).

Among various types of human neoplasms, the primary bone tumor is relatively uncommon. There is limited information regarding the prevalence and risk factors of primary bone tumors, which is associated with their features (3). Benign bone tumors are more prevalent than primary malignant ones. Therefore, benign tumors are apparently oftentimes underrated, as these mostly show no clinical symptoms and cannot be easily diagnosed. On the other hand, primary bone tumors are falsely known to be in the category of metastases, such as carcinoma, melanoma, or hematologic malignancies (plasmacytoma, etc.) (6). According to estimates, 2810 people were diagnosed with cancer (1,620 men and 1,190 women), and 1,490 people died from bone and joint cancer in 2011. About 0.2% of bone cancer malignances in the USA are sarcomas. In addition, age-adjusted incidence rate is 0.9 per 100,000 people each year. The most common histological subsets are chondrosarcoma (30% in men and 29% in women), osteosarcoma (16% in men and 17% in women), Ewing’s sarcoma (14% in both), and chordoma (8% in men and 5% in women) (7).

PRIMARY MALIGNANT BONE TUMORS

OSTEOSARCOMA

This type of bone tumor is a kind of mesenchymal tumor, which is highly malignant and the osteoid is produced by the tumor cells (8). It has the utmost prevalence of non-hematologic, primary, pe-

Figure 1. Osteosarcoma of the left distal femur
diatric malignancy, which is prevalent in children aged 10-25 years. This type of sarcoma may appear in any bone. However, it is the most frequent in juxta-epiphyseal areas of long bones with rapid growth (9). Malignant spindle cells, generating osteoid and undeveloped bone, are a histopathologic sign of high-grade intramedullary osteosarcoma (10). Other signs include a disorganized structure of the bone, having a fine lacey trabecular pattern, and irregular osteoid masses, regardless of its normal bone formation (11). The classic type of osteosarcoma has fibrous or chondroid appearance with small osteoid masses. According to cytogenetic examinations, there are many complex chromosomal abnormalities that make osteosarcoma different in each person. Osteosarcoma differs from other sarcomas, for example, Ewing’s sarcoma, synovial sarcoma, and alveolar rhabdomyosarcoma, in that it does not have any relationship with recurrent chromosomal rearrangements (12). There are different genetic alterations in osteosarcoma, which are shown by molecular analyses and include overexpressed oncogenes, including MDM2, and deactivation of p53 and retinoblastoma (Rb) tumor suppresser genes (13, 14). It has been proved that invasive surgery combined with multi-agent chemotherapy can increase the long-term survival by up to 60% (15). Significantly, studies have shown that patients who were only treated with chemotherapy had a minor survival rate of 20%. Thus, it can be argued that a high fraction of osteosarcoma-cells show resistance to chemotherapy (16). Several studies have been conducted on surgical and medical treatments for osteosarcoma, but the survival rate is still below 30 years, and 40% of the patients expire due to cancer (17, 18). Figure 1 shows an osteosarcoma of the left distal femur.

**Chondrosarcoma**

It is a malignant mesenchymal tumor, which is distinguished by different cells that produce a chondroid matrix. Contrary to osteosarcoma and Ewing’s sarcoma, this malignancy is most prevalent in adults aged ≥ 40 (19). Chondrosarcoma mostly happens in the shoulder, hip girdle, and pelvis. This sarcoma is diagnosed at a late stage and has a poor prognosis due to its deep lesions (20). The most prevalent symptom is pain in the lesion area. Histopathologically, it has a continuum from various hyalines, such as hypocellular chondroid lesions with low mitotic action, to high-grade pleomorphic chondrosarcomas possibly having slight chondroid (21). In-vitro and in-vivo studies that blocked hedgehog signaling demonstrated a reduction in the growth of chondrosarcoma. It seems that P53 and RB alterations are related to high-grade chondrosarcomas with 96% involvement (22 - 25). It has been

![Chondrosarcoma of the right distal tibia](image-url)
stated that cyclin dependent kinase 4 (CDK4) expression is associated with chondrosarcoma progression. On the other hand, in-vitro studies indicated that CDK4 could be knocked down by short hairpin RNAs, leading to diminished colony formation. There are studies suggesting an epigenetic constituent to evaluate chondrosarcoma pathogenicity (26, 27). Both chemotherapy and radiotherapy have no effects on chondrosarcomas. It has been shown that chemotherapy resistance is caused by the multidrug-resistance gene (MDR-1) P-glycoprotein (28). Therefore, adjuvant or neo-adjuvant chemotherapy is not suggested for conventional treatment of chondrosarcoma. The standard cure involves surgical resection with no adjunct irradiation or chemotherapy. Metallic endoprostheses, allografts or alloprosthetic composites are usually needed in this treatment because all bone sarcomas have skeletal reconstruction (29, 30). Figure 2 shows a chondrosarcoma of the right distal tibia.

**Ewing’s sarcoma**

This type of sarcoma is the second mostly prevalent primary malignant bone tumor in youngsters and teenagers, which is accompanied by small, round, and blue cells (31). Ewing’s sarcoma is most present in long tubular bones such as femur, tibia, fibula, pelvic girdle, and the ribs (32). Histopathological characteristics include small round blue uniform non-differentiated cells having little cytoplasm (33). Ewing’s tumors are positive in surface antigen CD99/MIC2 based on immunohistochemical studies. However, CD99 has no specificity to this sarcoma. CD99 may be present in lymphomas, leukemias, and rhabdomyosarcoma (34). Many patients with Ewing’s sarcoma have (11; 22) (q24; q12) chromosomal translocation, encoding the EWS/FLI oncoprotein (35, 36). One of the theories for the tumor origin is that Ewing’s sarcoma is a mesenchymal-derived tumor, implicating either neural ectoderm or mesenchymal stem cells (MSCs). For instance, neuroectodermal surface antigens are expressed on the Ewing’s cells, which supports the neuroectodermal origin (37). Some other studies explained that MSCs or progenitor cells derive from Ewing’s sarcoma. The cell of origin is still unknown, despite the fascinating available data (38).

A standard neo-adjuvant chemotherapy for the treatment of Ewing’s sarcoma includes vincristine, cyclophosphamide, and doxorubicin by alternation with etoposide and ifosfamide in combination with irradiation, operation, or both (39). Event-free survivability and general survivability are respectively 65% and 82% in patients having localized disease, and 25% and 39% in patients with recognizable metastasis. The latest studies suggest intensive chemotherapy within a more shortened interval, with the intensification of alkylating agents (40). Surgical approach involves the resection

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**Figure 3. Ewing’s sarcoma of the left pelvis**
of lesions in the appendicular skeleton and selective resection of lesions in the axial skeleton. The findings of surgery showed a local recurrence rate below 10% (41). Performing neo-adjuvant chemotherapy and, subsequently, surgical resection in which all tumor cells are completely removed provide a better structure for ambulation or the upper extremity functionality (42). Figure 3 shows a Ewing's sarcoma of the left pelvis.

**BENIGN PRIMARY BONE TUMORS**

**Osteochondroma**

Most of the benign bone tumors (30%) are cartilaginous tumors, which are found in tibia and femur. Osteochondroma is commonly seen in the metaphysis and diaphysis, and it mostly occurs in the above bones (43, 44). Solitary osteochondroma (exostosis) mostly occurs at the age of ≥40, while the autosomal and hereditary types of cancer are observed at younger ages with shortened extremities and deformities (45, 46). Thick periosteal reaction, endosteal scalloping, and cortical hook are the most reported features. Removing and cutting the tumor by surgery is the best treatment so far (47). Figure 4 shows an osteochondroma of the right distal femur.

**Giant cell bone tumor**

Giant cell tumors (GCT) comprise 20% of benign bone tumors and it mostly presents at ages 20 to 40 (48, 49). It is often presented in long bones, 50–65% in the area of the knee (50). GCTs are described as huge cells with osteoclast function, which are surrounded by spindle-like stromal and monocytic cells (51, 52). They are benign in 80% of cases. Recurrence after surgical resection is 20–50%, while 10% of cases may become malignant (53, 54). The first-line therapeutic option for this tumor is curettage, followed by bone cement filling. However, this method is highly prone to recurrence (55). Adjuvant treatment, such as zinc chloride, bisphosphonates, phenol, liquid nitrogen and alcohol, is used to reduce the recurrence (56). Wide excision and surgical methods are used for aggressive tumors.
(57). The latest suggestion for the treatment is the use of a chemotherapy drug called denosumab, which is a monoclonal antibody that inhibits the osteoclastic differentiation and activation of GCT by blocking osteoclasts (58). Figure 5 shows a giant cell tumor of the right proximal tibia.

Osteoblastoma

Another benign bone tumor is called osteoblastoma, which is rare and includes 14% of bone tumors (59, 60). It occurs at ages 20 to 40 (61). Axial skeleton with spinal lesions is the most common site of the tumor (about 33% of cases), but it may occur in other bones, too (62). The best choice for treatment is medical treatment. Radiotherapy and chemotherapy are the subsequent choices and surgical removal is the last one if the other methods fail. In a few cases, osteoblastoma progressed to osteosarcoma (63, 64). Figure 6 shows an osteoblastoma of the right humerus.

Osteoma

It is another outgrowth of membranous bones, which is benign and is most prevalent in the paranasal sinuses, skull and long bones (65). It may grow on homoplastic bones and heteroplastic or eutoplastic tissues (43). Osteoma affects osseous tissue, which comprises condensed bone with a well-defined border. It does not have irregular surface or satellite lesions (66). It is difficult to diagnose it if it shows no symptoms. Inflammatory response is considered as one of the underlying mechanisms because its incidence rate is increasing (67). Solitary osteoma is a risk factor for Gardner’s syndrome if there are multiple tumors (68). Surgical resection is suggested for symptomatic patients (69). Figure 7 shows an osteoma of the left femur.
CONCLUSION

A group of neoplasms, which is prevalent in young adults and children, is bone tumor. These tumors are diagnosed mostly by radiographic evaluations, but it is believed that CT and MRI can be used for detecting the local extent of the tumor. The treatment of benign bone tumors is done mainly in symptomatic patients, or those who are at the risk of pathological fracture and deformity. Surgical removal is the most common treatment for this problem. Active treatment is considered as the best treatment for GCT because there is a risk of malignancy. However, no consensus exists on standards of treatment for most of the benign tumors. Because of their low prevalence, malignant bone tumors are not easily classified and categorized for the sake of management. Bone malignancies are very important since they are found within the first years of life and, therefore, they can considerably affect the lives of patients and their families. The outcome and survivability of primary malignant bone tumors have improved recently due to both medical and surgical developments. Parallel to this, the molecular and cytogenetic representation have developed significantly, which, combined with light/electron microscopy and immunohistochemical methods, has a contribution to better understanding of these tumors.

Conflicts of interest statement

Authors declare there is no conflict of interests.
References


SAŽETAK

Tumori kostiju, uključujući benigne i maligne lezije, ne metastaziraju. Međutim, mogu se pojaviti u bilo kom delu skeleta. Najčešće se javljaju na distalnom delu femura i proksimalnom delu tibije. Većina benignih tumora zahvata hrskavice i zovu se osteohondromi. Na osnovu dosadašnjih izveštaja, benigni tumori kostiju češće se javljaju od primarnih malignih tumora. Maligni tumori kostiju pripadaju drugom tipu tumora kostiju i najčešće se javljaju u prvim godinama života. Iz tog razloga, značajno utiču na opšte stanje bolesnika i život njegove porodice. Ovi tumori obuhvataju osteosarkom, hondrosarkom i Juingov (Ewing) sarkom. U ovom radu prikazani su epidemiologija, karakteristike, i lečenje najvažnijih tipova benignih i malignih tumora kostiju. Ovi podaci biće korisni lekarima, kao i ostalim zdravstvenim radnicima, kako bi bolje razumeli tumore kostiju i njihovo lečenje.

Ključne reči: tumor kostiju, maligni tumor, benigni tumor