Case Report:
Clear Cell Sarcoma (Malignant Melanoma of Soft Part): A Rare Soft Tissue Sarcoma

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Clear cell sarcoma is a rare malignant neoplasm usually presenting as soft tissue mass in foot of young adult. Ultrastructural studies reveal neuroectodermal origin with melanocytic differentiation. Thus it is considered as malignant melanoma of soft parts. Though the prognosis is apparently better as compared to the conventional melanoma, local recurrences and metastases are frequently seen. Large tumor size and extent of necrosis are factors for poor prognosis. A case of clear cell sarcoma with classical presentation is reported.

Key words: Clear cell sarcoma, soft tissue sarcoma, malignant melanoma.

Soft tissue sarcomas are a large and heterogeneous group of neoplasms. Traditionally, these tumors have been classified according to histologic features. However, histomorphologic, immunohistochemical and experimental data suggest that most, if not all, sarcomas arise from primitive multipotential mesenchymal cells, which in the course of neoplastic transformation differentiate along one or more lines.

They account for approximately 0.1% of all cancers, but in children about 5% of tumors are soft tissue sarcomas. Clear cell sarcoma of tendons and aponeuroses is a rare and histologically distinctive malignant soft tissue tumor. Many authors believe that these tumors are neuroectodermal in origin and are derived from deep melanocytic cells associated with tendons and aponeuroses. The most common site is in the foot, often near the Achilles tendon or plantar aponeurosis.

Case Report
A 17 year old male presented with a swelling on the medial aspect of right ankle. It was there for six months causing occasional pain and steadily increasing in size. There were no other symptoms. Examination revealed a firm, circular 3x3cm, non-tender, mobile swelling placed between the right medial malleolus and the Achilles tendon. Skin over the swelling was normal and mobile. Ankle movements were within normal range. No popliteal or inguinal lymph nodes were palpable. X-rays of foot showed no bony lesion. Excision of the swelling was carried out. Per-operatively the swelling was grayish white, present below the tributaries of saphenous vein and the deep fascia. Posteriorly it was tethered to posterior tibial vessels. Swelling was separated all around; complete excision was done with adequate skin cover [Fig.1]. Microscopic section showed a cellular tumor consisting of sheets and groups of tumor cells with moderate nuclear pleomorphism, prominent nucleoli and high N/C ratio in a background of fibrocollagenous and vascular stroma. Scattered giant cells were noted with infrequent mitosis. Focally prominent intracellular golden brown melanin like pigment was noted. Features were consistent with malignant melanoma. The patient made an uneventful recovery. He is undergoing regular follow-up.

Figure 1: Post-operative photograph showing the area of lesion

Discussion
Clear cell sarcoma is a rare soft tissue neoplasm, arising most commonly in the tendons and aponeuroses of young adults. Though foot is the most common site, the neoplasm has been described in ankle, knee, thigh and other sites including penis. The largest study is of thirty-five cases of clear cell sarcoma, which were studied to determine the clinical or morphologic features that are important in predicting prognosis. Clear cell sarcoma usually presents as painless, gradually enlarging mass in young adult as seen in our case. Grossly the tumor is usually firm, well-circumscribed, grayish white and has a
gritty sensation on cutting. This sarcoma is characterized by small clusters of polygonal to spindle cells featuring clear to slightly basophilic cytoplasm and vesicular nuclei with prominent nucleoli. The clusters are separated by delicate fibrous septa. In a deletion, clear cell sarcoma has low mitotic activity, little or no necrosis and mild nuclear pleomorphism. Presence of multinucleated giant cells and abundant iron deposits have been described. Clear cell sarcoma of soft parts can be diagnosed readily by FNAC alone without resorting to incisional biopsy. In many cases the tumor cells contain cytoplasmic melanin. The diagnosis can be made with immunocytochemical confirmation of HMB-45 or S-100 protein expression, cytogenetic demonstration of the translation or electron microscopic studies demonstrating melanosomes. It is interesting that the same gene is also affected in Ewing’s sarcoma, PNET and intra-abdominal desmoplastic small-round cell tumor. The prognosis is relatively poor and local recurrences are quite frequent. Unlike most sarcomas, spread to regional lymph node is common, occurring in 30% cases. Metastatic spread has been reported in around 65% of patients and large tumor size along with degree of necrosis are predictors of poor prognosis. Complete surgical resection represents the main stay of treatment and possibly the only treatment for patients with small tumors. Radiotherapy may control microscopic residual disease after surgery. Chemotherapy is ineffective. The prognosis is unfavorable for non-resectable and large tumors.

References