Atypical Presentation of Ewing’s Sarcoma

Abstract

Background: Ewing’s sarcoma (ES) is an osseous malignancy of small round blue cells which may manifest even in soft tissue. It is the second most common primary tumor of bone in childhood. The common areas of occurrence are the diaphyses of femur followed by tibia, humerus, pelvis, and clavicle. However, there are many unusual presentations of ES on the basis of the site, e.g., craniofacial bones, paravertebral mass, and visceral organs. Chemotherapy with radiation therapy and cytoreductive surgery/limb salvage surgery is the mainstay treatment along with a strict patient compliance and counseling, critically being important for long-term survival. This study depicts presentations of ES who either presented with some usuality at the time of diagnosis or developed peculiar features uncharacteristic of ES during their course of treatment. Context: This study provides an insight into the patterns of unusual presentation of ES and prognosis of such patients with current line of management. Multiple sites of metastases have a very poor dismal outlook. Aims: This study purports the importance of considering ES as a systemic disease rather an osseous malignancy and reviews the pattern of unusual sites of presentation. Setting and Design: This observational study was carried out in the Department of Radiotherapy, Gandhi Medical College, Bhopal. Only patients with unusual patterns of metastases were included in the study cohort. Materials and Methods: We systematically reviewed patients with confirmed and immunohistochemistry-proven ES from July 2014 to July 2017. A total of 69 patients were registered within the time frame. Of the 69 patients, 36 (52.2%) were males and 33 (47.8%) were females. Of the 11 (16%) unusual presentations, 8 were males and 3 were females. The mean age of presentation was 22.5 (3–52 years). All patients received chemotherapy as per treatment guidelines. Each patient was individually followed up and metastatic workup was performed in a systematic manner. Statistical Analysis: Since the pattern of metastasis was observed, no significant statistical analysis was required for this study. Results: Of the 11 patients, six succumbed during the course of treatment, one patient was lost to follow-up, while four patients are on regular follow-up. Of the six patients who succumbed, four had multiple metastases at the time of presentation to our outpatient department, while the other two patients had paraspinal ES with paraplegia. Two patients, one male and one female, had associated comorbidities with spherocytosis and bronchial asthma. The youngest patient was a 3-year-old child with paraspinal ES, while the oldest was a 52-year-old with extraosseous ES of the right thigh who presented with lung, liver, and supraclavicular metastases. Conclusion: In our study, pattern of metastases determined the patients’ survival. Those patients who presented with multiple metastases at the time of presentation had a very poor prognosis, with death occurring within 2–3 months. The chemotherapy protocols were neither modified nor changed. They were evaluated after every cycle. Those patients who had developed multiple metastases at the time of admission to our outpatient clinic fared the worst. Those who had metastases to the brain and lung also succumbed to the malignancy. It is time we consider ES as an osseo-systemic malignancy and incorporate newer techniques such as circulating tumor cells in the investigation and evaluation portfolio for better and aggressive management. A multidisciplinary tumor board is absolutely essential and weekly meetings mandatory to individualize the treatment scenario. In our study, uncustomary as it may seem, their metastatic sites demand a certain degree of limelight in the Annals of Oncology.

Keywords: Age, Ewing’s family of tumors, metastases, unusual sites

Introduction

Ewing’s sarcoma (ES) was discovered in 1921 by James Ewing as “perivascular endothelial myeloma.”[1-2] The cell of origin of ES is not completely understood and consists of primitive neuroectodermal cells in varying stages of differentiation. ES is a malignant small round blue cell tumor in which the cancer cells are found in the bone or soft tissue. It is the second most common primary tumor of the bone in childhood. The most common areas of occurrence are the diaphyses of femur followed by tibia, humerus, pelvis, and clavicle.
clavicle. However, there are many patterns of unusual presentations of ES on the basis of the site of presentation, e.g., craniofacial bones, paravertebral mass, and visceral organs. The symptomatology depends on the primary site, but systemic symptoms with pain, fever, and weight loss are also seen. This study mainly focuses on the patterns of unusual spread in ES and provides an insight into the unwonted and unprecedented sites with which our patients symptomatically endured. Various literatures on ES focus mainly on rare sites such as craniofacial, hands and feet, gastrointestinal organs, and spinal masses. There have been case reports on ES on the ovary,[3] kidney,[4] and phalanges of hand.[5]

**Materials and Methods**

A total of 69 patients had been registered under our department with confirmed biopsy and immunohistochemistry (IHC)-proven ES from July 2014 to July 2017. Among them, 11 patients were seen to have unfamiliar patterns during presentation. They were marked for study and therapy initiated on the same lines of other ES patients. The patients involved in the study and their caregivers were informed through oral and written consent in their own language. Guarded prognosis was explained to the caregivers in view of the aggressive nature of the disease. Doses were calculated based on body surface area in adults and per kilogram in children. Of the 11 patients, eight were males and three were females. The age of patients ranged from a 3-year-old child to a 52-year-old male. The median age was 22.5 years. A detailed baseline and biochemical workup including lactate dehydrogenase levels were assessed pretherapeutically. Thorough history and physical examinations were performed.

**Results**

Of the 11 patients, four patients had ES, five extraosseous Ewing’s sarcoma (EES), one with primitive neuroectodermal tumor (PNET), and one with Askin’s tumor. Two had associated comorbidities, spherocytosis and bronchial asthma. The patient with Askin’s tumor developed scalp and leptomeningeal metastasis and succumbed. All 11 patients were IHC proven, and all of them were CD99 positive [Figures 1-2]. Five patients were vimentin positive and two were neuron-specific enolase positive along with CD99 (MIC-2). Five patients had multiple metastases at the time of presentations, of which four poorly responded to chemotherapy and succumbed very early during the course inspite of compliant treatment, while one patient still continues to follow-up. The two patients who had paraspinal ES developed acute-onset paraparesis and incontinence which progressed to paraplegia within a couple of weeks. Among them, the child was lost to treatment follow-up, while the other patient underwent debulking surgery and was initiated on adjuvant chemotherapy but died after 2 weeks. There were two patients who had a lag time of >2 years to develop metastasis who are still on regular review [Table 1]. Of the 11 patients, only four continue to review in our outpatient clinic while six succumbed and one patient was lost to treatment follow-up. Of the 11 patients, only one patient received radiation to primary site after six courses of chemotherapy, but yet after 3 months, he developed orbital metastases which were cytologically proven.
**Discussion**

ES is the third most common tumor in childhood with leukemia and CNS malignancies in the first and second place, respectively. Males are slightly affected more than females. The peak incidence in males is between 10 and 14 years of age, while in females, it is 5–9 years of age. The majority of patients seen for medical treatment are between 5 and 30 years of age. The most common sites in order of frequency are the pelvis, femur,ibia, fibula, rib, scapula, vertebra, and humerus. ES primarily afflicts the diaphyses of long-bone femur, humerus, clavicle, and pelvis.

Microscopically, they are sheets of uniform undifferentiated, small, deeply staining round blue cells. The differentials include lymphoma of bone, rhabdomyosarcoma, metastatic neuroblastoma, small-cell osteosarcoma, osteomyelitis, metastatic small-cell carcinoma of the lung, and mesenchymal chondrosarcoma. The characteristic immunostain is CD99 (MIC-2) which diffusely marks the cell membrane which is corroborated with chromosomal translocations, of which t(11;22)(q24;q12) is the most common present in about 90% of ES which codes for EWS and FL11, respectively, though there other gene defects of fewer penetrance example as ERG on Ch 21q22.3, ETV1 on Ch 7p 21.3, ETV4 on Ch 17q 21.31, EWSR1 on Ch 22q 12.2, FEV on Ch 2q 36, FLI1 on Ch 11q 24.1-q24.3, and FUS on Ch 16p 11.2.[6]

The Ewing’s family of tumors comprises the following:

1. ES
2. EES
3. Askin’s tumor
4. Primitive neuroectodermal tumor.

A multidisciplinary team comprising oncoseurgeon, radiation oncologist, medical oncologist, and onco-pathologist should concur with the treatment plan and chalk out an individualized management protocol.

Treatment basically involves chemotherapy, radiation therapy, and surgical resection, which depends on the localization of the lesion. First-line drugs used are vincristine, adriamycin, dactinomycin, and cyclophosphamide. Second-generation regimes with ifosfamide and etoposide have successfully improved the survival scenario in ES.

Alternating chemo schedule in the protracted format is the current treatment schedule given.[7,8] Radiotherapy has paid rich dividends in achieving local tumor control and pain palliation in patients treated with extensive bone lesions. Preoperative radiotherapy in doses 36–63 Gy has helped in reducing the size of extensive soft-tissue lesion in patients unfit for surgery after two cycles of chemotherapy.[9,10] However, for the metastatic patients, chemotherapy is the only salvageable approach while radiation may be given to alleviate pain or bleeding.[11] In our patients, multiple metastatic sites portrayed a dismal prognosis, with death occurring within 2–3 months [Figures 1-3]. ES is an osseous malignancy with variable soft-tissue component.

One patient had extensive extraosseous component of Ewing’s sarcoma with no bony deformity associated with spherocytosis [Figure 4]. Patients with craniofacial ES have a very poor prognosis as such lesions have extensive soft-tissue component which may invade brain and adjacent structure [Figure 5]. Any young patient presenting with acute neurological deficit with radiculopathy, paraplegia, and incontinence should be investigated with biopsy, and IHC and differentials must include ES. The oldest patient of the group was a 52-year-old male presented with an EES over the right thigh. He defaulted treatment for over 2 years and presented with multiple metastases in the liver, lungs, and supraclavicular lymph node. The unusual pattern in this patient is the age at which he presented. EES may present as soft-tissue swelling which may be located even intra-abdominally as seen in the patient who had PNET of the left kidney and the child who had anterior abdominal wall ES. Both the patients are compliant and still under...
follow-up. Patients with orbital metastases also have dismal outlook [Figure 6].

**Conclusion**

Our study shows that patterns of metastasis can determine the patients’ survival. Those who presented with multiple metastasis at the time of presentation to our department had a very dismal outlook with death occurring within the following 2-3 months. A multidisciplinary tumor board is mandatory to evaluate such patients and newer ancillary techniques to be incorporated in the investigation portfolio for aggressive management on an individual basis.

**Informed consent**

Prior written informed consent was taken from the patients and bystanders in their own language and either left thumbprint or signature was taken and collected.

**Declaration of patient consent**

The authors certify that they have obtained all appropriate patient consent forms. In the form, the patient has given his consent for his images and other clinical information to be reported in the journal. The patient understands that name and initials will not be published and due efforts will be made to conceal identity, but anonymity cannot be guaranteed.

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Nil.

**Conflicts of interest**

There are no conflicts of interest.

**References**

4. Hakky TS, Gonzalvo AA, Lockhart JL, Rodriguez AR. Primary Ewing sarcoma of the kidney: A symptomatic presentation and