Post Traumatic Dorsal Mass of a Child Hand: Rhabdomyosarcoma Must Be Suspected

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Abstract

Rhabdomyosarcoma (RMS) is a highly malignant soft tissue tumor accruing in children. Strongly suggestive history of trauma is recognized as an etiological factor of RMS. Rapidly growing, persistent firm, or non-mobile lesions appearing after trauma should raise the suspicion of malignancy. If doubt exists, radiological exams must be performed because appearance of RMS is highly suggestive. We report a case of 8-year-old girl with a history of trauma how’s developed RMS of the dorsum of his left hand and we describe radiological appearances suggesting this diagnosis.

Keywords: Rhabdomyosarcoma (RMS); Soft tissue tumor

Introduction

Peritoneal Rhabdomyosarcoma (RMS) is the most frequent, highly malignant soft tissue tumor that develops from striated muscle cells [1]. History of trauma is recognized as an etiological factor of RMS [2]. But trauma and a small painless mass is apt to be overlooked or misdiagnosed unless the physician is constantly aware of the possibility of this possibility of malignant evolution. We report a case of 8-year-old girl with a history of trauma how’s developed RMS of the dorsum of his left hand and we describe radiological appearances suggesting this diagnosis.

Case Report

An 8-year-old white child without medical history was brought for a swelling of the dorsum of her left hand. Her mother states that the mass appeared almost one year ago, few days after the girl strucked her hand when she fell, and has gradually increased in size despite a medical symptomatic care.

On physical examination, the patient weighted 20 kg (-3 Standard deviation), she was asthenic and has increased temperature of 38°C.

The mass has ill-defined limits, was roughly rounded and measured 4 cm × 5 cm. It was firm, painful on palpation, and fixed relative to the deep plane of the dorsal 4th and 5th metacarpals.

Another painful mass was found in the inner side of the ipsilateral elbow which was 3 cm in long. An axillary swelling lymph node was discovered too.

Biological exams showed pancytopenia, elevated sedimentation rate, hypercalcaemia and ascent of muscle enzymes.

Antéro posterior X-ray of the hand and elbow (Figure 1) revealed a soft tissue swelling over the dorsal metacarpals and an irregular erosion of the second right metacarpal. Strangely, another erosion was seen in the lower extremity of the ulna.

An ultrasound of soft tissues of the hand was performed the lesion was heterogeneous hypo echoic highly vascularized in color Doppler, surrounding the 3rd and 4th and 5th metacarpals (Figure 2). At magnetic resonance imaging, the mass arises from the dorsal and palmar interossei and occupied both the dorsal and palmar compartments of the hand. The mass was heterogeneous hypointense to the muscles on T1WI and T2WI. Marked homogeneous enhancement was demonstrated after gadolinium injection (Figure 3).

These findings suggested a malignant soft tissue mass that most likely originated from the interossei muscle and infiltrated the third metacarpal. Spinal MRI showed multiple vertebral collapses with heterogenous spinal matrix suggesting bone marrow involvement. Chest CT scan was performed and revealed three parenchymal metastatic nodules.

A metastatic RMS was diagnosed. Excision biopsy of the lesion was performed through a dorsal approach. The histology of the specimen was consistent with RMS showing alveolar subtype. The postoperative period was uneventful. And the patient received four intensive treatments.

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courses of combination therapy using Vincristine, Adriamycin and Cyclophosphamide (VAC), together with radiotherapy.

Discussion

RMS is a highly malignant soft tissue tumor that develops from striated muscle cells. It accounts approximately 55% of all soft tissue sarcoma in children younger than 15 years and is more common in boys [1,2]. Pathogenesis of the tumor is uncertain. Strongly suggestive history of trauma is recognized as an etiological factor of RMS [3]. The trauma can occur years before the appearance of the tumor or a very short time before its recognition. In both cases, rapidly growing, persistent firm, non-mobile lesions should raise the suspicion of malignancy [4]. Also, biological disturbance as in our described case should have drawn the attention and suggested malignancy. If doubt exists, radiological exams must be performed because appearance of RMS is highly suggestive. The couple radiography and ultrasound should be the two initial imaging studies of a suspicious hand mass as radiography and ultrasound can offer highly sensibility detecting little calcifications described in 10% to 30% of hand RMS [1]. MRI which provides high soft tissue and spatial resolution is essential to evaluate the extent of the mass. A bone scan also should be part of the preoperative evaluation. Computerized tomography (CT) results are more informative than MRI results when evaluating cortical changes and assessing calcification within the lesion [1,7]. Extremity RMS is characterized by its particularly poor outcome and its high tendency to develop distant metastases. So that 5-year survival RMS of hand and limb is of 33.3% [8] and 29% of patients had distant metastases at the time of diagnosis while the usual rate for RMS of all sites is 10% to 20% [8,9]. Chest CT scan should be performed to detect metastatic disease [5]. This exam showed three metastatic nodules in the case of our patient. In opposition to bone tumors, soft tissue tumors, especially RMS can metastasize to regional lymph nodes [9,10]. In our patient, the second painful mass of the inner side of the ipsilateral elbow corresponded surgically to a metastatic lymph node. Axillary lymph node was also resected and was metastatic. Another important characteristic is the high tendency of RMS to develop bone marrow dissemination.

Conclusion

Our reported case confirms this concept since our girl has bown marrow invasion at the time of diagnosis attested by pancytopenia, vertebral collapse and heterogenous spinal matrix on spinal MRI. In addition, even if no biological abnormality is found and because of the highest tendency of metastatic RMS to involve bone marrow, bone marrow aspiration should be a part of pretherapeutic evaluation of such children [2,10]. Histologically, RMS has been assigned to three histological categories: embryonal, alveolar and pleomorphic with 15% to 20% of rhabdomyosarcomas not fitting into a conventional classification and remaining either undifferentiated or unclassifiable tumors [5]. Alveolar subtype found in our case is the most frequent RMS of the extremities and has the worst prognosis [11]. Radical surgical ablation by en-bloc excision or amputation affords the best chance of survival [2,12]. The use of multiple agent chemotherapy regimens (for example, VAC) produced an immediate improvement in survival in patients with lesions in the extremities.

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