Osteomyelitis of Myositis Ossificans in Arm – First Case Report

Pankaj Kumar Mishra¹, Prakhar Singhal¹, Jiten Shukla¹, Deepak Singh Maravi¹

What to Learn from this Article?
A very rare disease presentation and management

Abstract

Introduction: Myositis ossificans [MO] is a benign heterotrophic bone forming (often self-resolving) pathology of bone and soft tissue. Here we are reporting the first time in literature for osteomyelitis of myositis ossificans in arm of a male due to trauma as a perusal of rare entity.

Case Report: It is a case report of a 25 years old male presented to us in out-patient department with chief complaint of discharging wounds over mid part of left arm since six months. Clinically provisional diagnosis of chronic osteomyelitis of left humerus made and his x-ray sought. X-Ray showed geographic appearance of myositis ossificans around the upper two third of left arm. Sinuses curetted and infected bone (part of myositis ossificans) removed and sent for biopsy. Now the patient is discharge and sinus free, and has resumed his work.

Conclusion: Osteomyelitis of myositis ossificans should be recognized as a possible differential diagnosis chronic discharging sinus. This type of presentation of myositis ossificans is rarest.

Keywords: Osteomyelitis, myositis ossificans, arm

Introduction

Myositis ossificans [MO] is a benign heterotrophic bone forming (often self-resolving) pathology of bone and soft tissue. It has also been reported to occur due to strain or overuse of muscles. Here we are reporting the first time in literature for osteomyelitis of myositis ossificans in the arm of a male due to trauma as a perusal of a rare entity.

Case Report

It is a case report of a 25 year old male presented to us in out-patient departments with chief complaint of discharging wounds over mid part of left arm since six months. On the complete history taking he revealed that he had an episode of trauma one year back over his left arm due to fall on the ground and swelling develops at that time. For it he did not take any medical advice and he took few pain killers and repeated...
Mishra PK et al

Discussion

The exact pathogenesis of MO is not clear but one theory stated that due to injury there are fibroblastic proliferation or osteoblastic migration (from injured periosteum) into haematoma, have been blamed for causing the pathology of myositis ossificans [2]. Neither is it found exclusively in skeletal muscle nor there is muscle inflammation so its name is misnomer, even the name myositis ossificans is most commonly used clinical term. So the term heterotrophic ossification seems clinically worth [3,4]. Myositis ossificans have three variants, myositis ossificans circumscripta (MOC), neurogenic myositis ossificans and fibrodysplasia (myositis) ossificans progressiva.

Myositis ossificans circumscripta (or traumatica) is a pseudosarcomatous pathology with restricted growth tendency. If its excision is done in matured phase (well demarcated from surrounding tissue) and if repeated injury is avoided then it may prevent the recurrence of it [5].

Neurogenic form of myositis involves the large joints of the body (hip, knee) if they have been immobilized due to traumatic neurological damage, burn or arthroplasty. But on the contrary to circumscripta, it involves the connective tissue between the muscle plane and around the joint rather than skeletal muscle [6, 7].

Fibrodysplasia (myositis) ossificans progressiva (FOP) have the preponderance for young age while the other two forms do so rarely. FOP is an autosomal dominating inherited disabling disease (a disorder of bone morphogenetic protein 4) have a predilection to paraspinal, scalp and temporomandibular joints (but never involves facial muscles, tongue, diaphragm and viscera). FOP has the characteristic feature of (potentially recognizable at birth) short big toe, hallux valgus (also thumb and cervical spine sometimes involved) and broad femoral neck. There is spontaneous (also by trauma) enchondral ectopic ossification in tendons, ligaments, joint capsule and progressively in a specific manner and give a feature of “second skeleton” [8, 9, 10].

MOC has peculiar clinico-radiological and cytological feature and clinically present as painful soft tissue swelling with restricted range of movement in a joint following a trauma. In the early stage it appears as irregular, hazy, flocculent structure, but after 3 weeks zonal calcified area is evident. So for the early diagnosis of MOC, sonography and three phase bone scan is needed [11, 12]. This characteristic radiological feature of “zoning,” is due to distinct mature ossification at the periphery and centrally situated radiolucent (which is due to immature osteoid and primitive mesenchymal tissue) nidus [5]. This characteristic feature of “zoning” differentiate it from extra skeletal osteosarcoma in which there is centrally situated radio opaque matured osteoid cell. Zonal architecture is obvious after 6 weeks of trauma and Computer tomography is needed to demonstrate this zonal mineralization pattern [13].

For the management of MOC initially conservative management (i.e. - rest, immobilization and physiotherapy) should be given and surgical excision can be used as a last resort for the matured stage (6-12 months of trauma) to avoid recurrence. Even after the diligent search we did not get any article on osteomyelitis in MO. Etiopathogenesis of osteomyelitis in myositis mass can be conceded same, as it occurs in other bone (i.e. direct inoculation due to trauma or hematogenous spread). In our case the osteomyelitis would have been occurred via hematogenous spread. Our case had the feature of chronic osteomyelitis (chronic discharging sinuses and protruded sequestrum) so the treatment protocol was the same as usual of the chronic osteomyelitis of normal bone.

In summary, as a rare case in literature we are presenting the case of osteomyelitis of MOC due to trauma. In this case MO was in matured stage and infected so the surgical excision done. As a perusal of rare presentation, clinicians should be aware of this unusual presentation of MO.

Conclusion

Osteomyelitis of myositis ossificans can be conceded as a possible differential diagnosis of chronic discharging sinus in a few feasible conditions (following trauma around joints). This type of presentation of myositis ossificans is rare, and we are...
reporting this is as a first time in literature.

Clinical Message
This unusual presentation of myositis ossificans is not mentioned in literature, in our knowledge. So we are bringing it to horizon of knowledge to disclose the unusual presentation of myositis ossificans.

References

Conflict of Interest: Nil
Source of Support: None

How to Cite this Article