Clinical Presentation and Management of Trigonum Bone Syndrome in a Pediatric Patient

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Abstract

This article presents a rare case of Trigonum Bone Syndrome in an 8-year-old girl, diagnosed using X-ray, Computed Tomography (CT) and contrast-enhanced magnetic resonance imaging (MRI). The patient presented persistent ankle pain and ligamentous distention, not responding to conservative treatments. Surgical removal of the accessory bone and subsequent rehabilitation led to successful outcomes.

Keywords: Talus; Rare diseases; Ankle joint; Tarsal Bones; Trigonum Bone Syndrome; Os Trigonum.
MRI provided the evidences of adjacent soft tissue edema and inflammation, suggesting the presence of Trigonum Bone Syndrome (Figure 3). There was no evidence of osteochondral lesions or other bony abnormalities.

After discussing the available treatment options with the patient's parents, the decision was made to proceed with surgical intervention. The patient was admitted for an excision of the os trigonum and exploration of the surrounding soft tissue structures. The surgery was performed under general anesthesia, with the patient in a prone position. A posterolateral approach to the ankle joint was utilized, with careful dissection of the soft tissues to protect the neurovascular structures. The os trigonum was identified, and its fibrous attachments to the talus were released. The accessory bone was then excised, and the surrounding soft tissues were carefully inspected for any signs of injury or inflammation. The wound was closed in layers, and a sterile dressing was applied.

Postoperative recovery was uneventful, with the patient experiencing significant pain relief and improved range of motion in her right ankle. She was referred for rehabilitation, which included a gradual return to weight-bearing activities and a tailored physiotherapy program to strengthen the surrounding musculature and improve joint stability.

Discussion

Trigonum Bone Syndrome is a rare condition that occurs due to the presence of an accessory bone, the os trigonum, in the talus. The presence of this extra bone can lead to discomfort, pain, inflammation, and limited mobility in the ankle joint that is affected. Although the presence of an os trigonum is relatively common, symptomatic cases are rare and often require surgical intervention [5].

The etiology of Trigonum Bone Syndrome is thought to be congenital, arising from the non-fusion of an accessory ossification center during embryological development [1]. The ossification center for the os trigonum appears around the 10th week of gestation and typically fuses with the primary ossification center by the age of 8 to 13 years [2]. Failure of this fusion process results in the presence of an os trigonum.

Diagnosis of Trigonum Bone Syndrome can be challenging due to its rarity and non-specific clinical presentation. In this case, the initial diagnosis was made using plain radiographs, which showed the presence of an accessory bone in the talus. Further evaluation using a CT and a contrast-enhanced MRI helped to confirm the diagnosis and assess the extent of soft tis-
sue involvement [6]. The findings of Magnetic Resonance (MR) imaging include edema in the talus and/or os trigonum, fluid in the synchondrosis, fluid surrounding the os trigonum, and alterations in soft tissue. Additionally, MR can reveal related findings such as stenosing tenosynovitis of the flexor hallucis longus and degenerative changes around the synchondrosis or between the os trigonum and the adjacent calcaneus.

Treatment options for Trigonum Bone Syndrome vary depending on the severity of the symptoms and the extent of soft tissue involvement. Conservative measures, such as rest, ice, compression, elevation, and analgesics, are often recommended initially [7]. In cases where these measures fail to provide adequate relief, as in our patient, surgical intervention may be necessary.

Several surgical approaches have been described for the excision of the os trigonum, including open and arthroscopic techniques [8]. In this case, an open posterolateral approach was utilized, which allowed for direct visualization of the os trigonum and the surrounding soft tissue structures. The choice of surgical approach should be based on the surgeon’s familiarity with the technique, the patient’s anatomy, and the extent of soft tissue involvement.

Postoperative management typically involves a period of immobilization followed by a gradual return to weight-bearing activities and a tailored physiotherapy program. In our patient, the postoperative recovery was uneventful, with significant improvement in pain and range of motion in the right ankle. This highlights the importance of early diagnosis and appropriate surgical intervention in achieving successful outcomes in patients with Trigonum Bone Syndrome.

In light of the rarity of Trigonum Bone Syndrome, increased awareness among clinicians and surgeons is essential. Early diagnosis and appropriate management can significantly improve patient outcomes and prevent potential complications, such as chronic pain, instability, or reduced range of motion. Furthermore, understanding the embryological basis of this condition can contribute to the development of more targeted and effective treatments in the future.

It is important to note that while our case report focuses on a pediatric patient, Trigonum Bone Syndrome can affect individuals of all ages. In particular, it has been reported in athletes who engage in repetitive plantarflexion activities, such as ballet dancers and soccer players [4,3]. In these populations, early recognition and treatment are crucial to avoid potential long-term consequences and to facilitate a safe and timely return to sports.

Additionally, clinicians should be aware of the potential differential diagnoses when encountering a patient with posterior ankle pain and an accessory bone in the talus. These may include conditions such as an ununited fracture fragment, osteochondroma, or a Stieda process [5]. Careful evaluation of the patient’s history, clinical presentation, and imaging findings can help to differentiate these entities and guide appropriate management.

Lastly, this case report highlights the role of advanced imaging techniques, such as CT and contrast-enhanced MRI, in the diagnosis and management of Trigonum Bone Syndrome. As the condition is characterized by the involvement of both osseous and soft tissue structures, the use of MRI can provide valuable information on the extent of the pathology and aid in surgical planning [3].

Trigonum Bone Syndrome is an unusual but important cause of posterior ankle pain, particularly in pediatric patients and athletes. Early diagnosis and appropriate management, including surgical intervention when necessary, can lead to successful outcomes and prevent long-term complications. Clinicians and surgeons should maintain a high index of suspicion for this condition and be familiar with its diagnostic and treatment options. Further research is needed to improve our understanding of the etiology, natural history, and optimal management strategies for this rare condition.

**Conclusion**

Trigonum Bone Syndrome is an uncommon condition identified by the existence of an accessory bone within the talus. This can cause discomfort, pain, inflammation, and diminished mobility in the ankle joint that is affected. The diagnosis can be challenging due to its rarity and non-specific clinical presentation, but imaging studies such as X-ray, CT and contrast-enhanced MRI can aid in the diagnostic process. Surgical intervention is often necessary in cases where conservative measures fail to provide adequate relief. In this case, surgical excision of the os trigonum and subsequent rehabilitation led to a successful outcome, highlighting the importance of early diagnosis and appropriate management in patients with this rare condition.

**Declarations**

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**References**