Chronic Active Parietal Osteomyelitis Due to *Salmonella typhi* in a Patient with Sickle Cell Anemia

Orak Hücreli Anemi Hastasında *Salmonella typhi* Kaynaklı Kronik Aktif Parietal Osteomiyelit

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To the Editor,

Sickle cell disease (SCD) is a genetic disorder characterized by marked heterogeneity in clinical and hematologic severity, with musculoskeletal system manifestations being a major cause of morbidity and disability [1]. The increased susceptibility of SCD patients to infections, including osteomyelitis, has long been recognized with several mechanisms postulated including impaired splenic function, defects in complement activation, genetic factors, deficiencies in micronutrients, and the presence of infarcted or necrotic bone [2]. *Salmonella* is the most common cause of osteomyelitis in SCD, followed by *Staphylococcus aureus* and gram-negative enteric bacilli; this prevalence could be related to the fact that intravascular sickling of the bowel leads to patchy ischemic infarction [3,4]. The most common sites of osteomyelitis are the femur, tibia, or humerus. Patients usually present with acute onset of pain, swelling, and tenderness over the affected area in association with fever and elevated inflammatory markers. However, in some cases, osteomyelitis has atypical presentations with a more indolent course and often with abscess formation [5]. Here we present a 50-year-old female patient with sickle cell anemia (SCA) who developed parietal osteomyelitis with abscess formation and involvement of the dura due to *Salmonella typhi*, who was treated successfully by surgery followed by antibiotics.

A 50-year-old Saudi female patient living in the Eastern Province of Saudi Arabia, diagnosed with SCA (HgS: 78%) with occasional vaso-occlusive crisis and no sickle cell-related complications, presented to us with a 1-month history of a painless right parietal subgaleal collection increasing in size over time with no history of trauma and no fever or neurological manifestations. Laboratory testing revealed an elevated white blood cell count and a high estimated sedimentation rate level (125 mm/h). Magnetic resonance imaging of the brain revealed an osteolytic defect centered on the right parietal bone and sizable subgaleal complex collection (Figure 1). The patient underwent right parietal craniectomy with cranioplasty (removal of the right parietal subgaleal collection and the corresponding bone in addition to the invaded dura). Pathology of the specimen revealed a right parietal subgaleal abscess and right parietal bone chronic active osteomyelitis. Culture of the specimen grew *Salmonella typhi*.

The morbidity of chronic osteomyelitis combined with other complications of SCD decreases patients’ quality of life. Patients with SCD are more prone to osteomyelitis. The most common causative organism is *Salmonella*. The usual manifestations of osteomyelitis are pain, swelling, tenderness, and fever. However, like in our case, sometimes osteomyelitis presents late, as a more indolent process often with abscess formation and in unusual and more critical sites. Our case highlights the atypical presentation of osteomyelitis in a patient with SCD, which could cause devastating complications if not treated properly, early, and by a multidisciplinary team approach.
Acquired Leukocyte Inclusion Bodies Resembling Döhle Bodies During Acute Cholangitis

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To the Editor,

A 66-year-old woman was admitted to the gastroenterology department with epigastric pain, nausea, and subicterus. Her complaints had begun 6 h earlier. Her abdomen was soft and flat, with localized tenderness on palpation in the right subcostal area. Laboratory studies revealed a white cell count of 17.9x10^9/L, hemoglobin concentration of 14.4 g/dL, and platelet count of 48x10^9/L, and they were notable for elevated serum cholestatic enzymes. The abdominal ultrasound was remarkable for cholangitis. The patient received broad-spectrum antibiotics. A peripheral blood smear examination, performed to evaluate thrombocytopenia, revealed the presence of blue intracytoplasmic inclusions in neutrophils (Figures 1A-C). On the 11th day of treatment, her blood smear was examined once again and the Döhle body-like inclusions were resolved (Figure 1D).

May-Hegglin anomaly is an uncommon autosomal dominant abnormality characterized by large, basophilic inclusion bodies (resembling Döhle bodies) in neutrophils [1,2]. Döhle bodies can be seen in bacterial infections. Hematologic findings of systemic diseases may be confused with hematological diseases such as May-Hegglin anomaly. We thought that the granules were Döhle bodies due to cholangitis. The disappearance of the inclusion bodies upon treatment is important in differential diagnosis.