Abstract

Sickle cell Disease is the most common genetic disorder in children living in sub-Saharan Africa [1]. The burden of disease is estimated to range between 2-4% in malaria endemic regions [2]. The disease is known to cause a number of acute episodes in patient presenting with SS, SC and S-beta-thal and also now known to cause multiple organ pathologies and long term morbidities and eventual mortality [3]. The pathologies commonly affect the brain (stroke), eyes (retinopathy), kidney (micro and macro albuminuria), Bones (acute pain, aseptic necrosis, osteomyelitis and skeletal deformities) heart (cardiomyopathy and pulmonary hypertension), Jungs (Acute Chest Syndrome) and overwhelming bacterial infections [3-9]. Early detection and appropriate follow up with prompt therapy and preventive measure like folic, penicillin V prophylaxis and immunization against encapsulated organisms like streptococcus pneumonia and Haemophilus Influenzae bacterial are known to improve survival and quality of life [10]. SCD presents with other comorbidities that complicate the disease process, resulting in diagnostic and management challenges. This case study presents a scenario of a sickle cell patient with multiple long term complications and Deep Vein Thrombosis (DVT) as a co-morbid condition. He did not benefit from early diagnosis and preventive management therapies. This 12-year-old known SCD patient presented with 2 months history of inability to walk, weight loss and pain in both legs to a tertiary hospital in Ghana. On examination he was wasted (Weight for Height z-score <-3SD) and pale. He had palpable axillary and inguinal lymph nodes. His right leg was swollen, warm, erythematous and tender. He had a gibbus deformity of the lower thoracic spine. Investigations revealed Hemoglobin 7.1g/dl, WBC of 13.51 X 109/l with Neutrophils (65.3%). ESR-117, INR-1.05, Prothrombin Time-14.3secs and blood sugar-6.5mmol/L. Radiological findings revealed features suggestive of sickle cell disease with Thoracolumbar spondylodiscitis, likely pyogenic. Bilateral Anasular Necrosis of the femoral head, Septic arthritis of the right hip and left shoulder joint, Osteomyelitis of the left 8th and 9th ribs. Doppler Ultrasound of the lower limb revealed Right Popliteal and Posterior Tibial Vein Thrombosis with Pyomyositis. The patient was treated with Enoxaparin, Warfarin and IV antibiotics (Clindamycin and Ciprofloxacin). Follow ups with a repeat Doppler finding revealed a completely resolved thrombus with no evidence of DVT and normal INR and PT. Attention therefore needs to be paid to a multi-disciplinary preparedness and response approach to the management of SCD patients, as presented in this case report, to improve the quality of life, mitigate disability and reduce mortality.

Introduction

Sickle cell Disease is the most common genetic disorder in children living in sub-Saharan Africa [1]. The burden of disease is estimated to range between 2-4% in malaria endemic regions [2]. The disease is known to cause a number of acute episodes in patient presenting with SS, SC and S-beta-thal and also now known to cause multiple organ pathologies and long term morbidities and eventual mortality [3]. The pathologies commonly affect the brain (stroke), eyes (retinopathy), kidney (micro and macro albuminuria), Bones (acute pain, aseptic necrosis, osteomyelitis and skeletal deformities) heart (cardiomyopathy and pulmonary hypertension), Jungs (Acute Chest Syndrome) and overwhelming bacterial infections [3-9]. Early detection and appropriate follow up with prompt therapy and preventive measure like folic, penicillin V prophylaxis and immunization against encapsulated organisms like streptococcus pneumonia and Haemophilus Influenzae bacterial are known to improve survival and quality of health [10]. SCD presents with other comorbidities that complicate the disease and presents with diagnostic and management challenges. This case study presents a scenario of a sickle cell patient with long term complications and Deep Vein Thrombosis (DVT) as a co-morbid condition, who did not benefit from early diagnosis and preventive management therapies.

Case Report

A 12-year-old boy was diagnosed with Sickle Cell Disease (SCD) with genotype SS about one year prior to. He presented with 2 months history of inability to walk and pain in both legs to Komfo Anokye Teaching Hospital in Kumasi, Ghana. The complaints were associated with marked weight loss and generalized body pain. He had been admitted about 2 weeks prior to the onset of current symptoms on account of abdominal pain and anaemia where he was haemo-transfused, given medications and subsequently discharged. There were no other significant findings in the past medical history, immunization history, nutritional and developmental histories. On examination he was found to be wasted (Weight for Height z-score <-3SD), pale, afebrile and anicteric. He had palpable axillary and inguinal lymph nodes (each measuring about 1cm to 1.5cm, mobile, non-tender and not matted). Vital signs were normal on admission. Other significant examination findings were obvious swelling of the right leg which was warm to touch, erythematous and tender on palpation. He had a gibbus deformity of the lower thoracic spine with no remarkable neurological findings in the lower limbs. A diagnosis of SCD with vaso-occlusive crises and cellulitis of the right leg was made with differential diagnosis of osteomyelitis, Deep Vein Thrombosis and Potts disease. Initial investigations revealed WBC of 13.51 X 109/
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dl with Neutrophils 8.83 (65.3%) Lymphocytes 4.27 (31.6%) and Monocytes 0.25 (1.9%). Hemoglobin 7.1g/dL, MCV-70.7, MCH-22.4, Platelets: 538, ESR-117, INR-1.05, Prothrombin Time-14.3secs and blood sugar-6.5mmol/L. Malaria parasites were not seen on the blood smear and patient was negative for Retro-Viral Screen for HIV. Other investigations included Doppler Ultrasound, Chest X-ray, Pelvic x-ray.Thoraco-lumbo-sacral spine X-ray (Figure 1-10). Blood culture and sensitivity could not be done at the time of admission as microbiology service in the hospital had a brief logistic challenge. Radiological findings revealed the following: Reduction in the vertebral body heights of T7-T10 with associated reduction in the intervertebral disc spaces and crowding of the posterior ribs. A paravertebral soft tissue mass extending from T4 vertebral level to T11. Coarse trabeculation and sclerosis of the ribs noted. Expansion of the anterior end of the left 8th rib with ill-defined lytic areas. Buckling of the cortex of the lateral aspect of the left 9th rib, suggestive of a fracture. The imaged portion of the proximal humerus shows periarticular osteopenia. Also noted was relatively small left humeral head with left glenohumeral joint space. There are patchy homogenous opacities in both lung fields predominantly in the lower zones. There was associated silhouetting of the hemi diaphragms bilaterally and the cardiac shadow. There was generalized sclerosis of the imaged pelvic bones with coarsening of the trabeculae. The right femoral head was deformed and expanded with sclerosis and lytic areas within. There was associated narrowing of the right hip joint space. The left femoral head also showed patchy luencies. No deformity of the left femoral head seen. The left hip joint space appeared normal. Also noted was fusion of the sacroiliac joints bilaterally. There was diffuse sclerosis of the imaged vertebral with coarse trabeculae. The vertebral endplates of L1/L2 and L3/L4 showed sharp depression at the margins with a flat base centrally giving it the H shaped appearance. Also noted were ill-defined lytic areas in the L3 to L5 vertebral with associated end plate irregularities and reduced intervertebral disc spaces at L3/L4 and L4/L5. No paravertebral soft tissue mass seen. No fracture or listhesis seen. Changes of the thoracic spine were described under the chest x-ray.

These radiological diagnostic features were suggestive of sickle cell disease with Thoracolumbar spondylodiscitis, likely pyogenic; Bilateral Avascular Necrosis of the femoral head (early on the left); Septic arthritis of the right hip and left shoulder joint; Osteomyelitis of the left 8th and 9th ribs. Doppler Ultrasound of the lower limb revealed the following: The right common femoral and superficial femoral veins were of normal lumen with good wall to wall compressibility, good colour doppler and spectral wave form properties. The right popliteal and posterior tibial veins however showed extensive echogenic intraluminal-filling defect with resultant loss of wall-to-wall compressibility (percentage stenosis 76%). The accompanying arteries were of normal size and caliber and showed good colour doppler and spectral properties. There was thickening of the subcutaneous tissue with dilated fluid channels seen in the right foot. Inguinal as well as popliteal lymph nodes with retention of their fatty hilum noted averaging 0.9*0.5cm. Also noted was a 1.4*0.9*1.5cm (volume about 1.0ml) anechoic collection with low-level internal echoes at the medial aspect of the proximal thigh muscles; suggestive of inflammatory changes. Based on the Doppler findings a clinical diagnosis of Right Popliteal and Posterior Tibial Vein Thrombosis with Pyomyositis in a Sickle Cell Disease patient was considered and patient was subsequently administered subcutaneous Enoxaparin 40mg daily as well as Warfarin 3mg nocte, IV Clindamycin 150mg qid, IV Ciprofloxacin. After 2 days of Warfarin and Enoxaparin, INR and PT were repeated and results were 1.1 and 15.1s respectively. The child was followed up closely for four weeks and subsequently discharged. A repeat Doppler finding revealed a completely resolved thrombus with no evidence of DVT.
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Figure 3: Pelvic X-ray of patient.

Figure 4: Lateral Lumbo Sacral spine.

Figure 5: AP view Lumbo Sacral spine.

Figure 6: Doppler Ultrasound of patient.

Figure 7: Doppler Ultrasound of patient.

Figure 8: Doppler Ultrasound of patient.

Discussion

Sickle cell disease a genetic disorder of the red cell but frequently associated with multiple end-organ complications if not diagnosed at birth and managed appropriately [11,12]. The patient presented in this case report showed up with variety of complications that can occur in SCD patients if adequate effort is not placed on early diagnosis and prevention of complications[11,13]. The history suggests that patient was not offered early diagnosis and initiation of preventive measure like the administration of Folic acid and Penicillin V. The occurrence of multiple pathologies related to SCD is therefore not unexpected. The level of wasting and malnutrition could not be attributed to the SCD alone since the existence of chronic diseases like tuberculosis or recurrent acute episodic infections in Sickle Cell Disease could contribute to the level of muscle wasting and malnutrition. Studies have shown that SCD patients are more likely to become malnourished compared to their age and sex match [14,15]. The lack of early diagnosis and management of the disease and the morbidities acquired are sufficient to cause wasting and malnutrition. It is well established that SCD patients are at risk of infections more than the general population [16-18]. The risk is estimated to be about 36 times for encapsulated infections [17]. *Streptococcus pneumonia*, *Haemophilus Influenzae Type b (Hib)* and *Salmonella Typhi and Para Typhi* are the most common organisms responsible for infection in SCD patient [17,19-21]. The occurrence of multiple bone and joint infections makes *Salmonella* species the most likely organism to be responsible for the septic arthritis of the joints and osteomyelitis. Another potential organism to cause such a disseminated infection in the patient is *Staphylococcus aureus* which together with *Salmonella* species are said to be the two most common causes of osteomyelitis in children with sickle cell anemia [22]. The occurrence of gibbus in a malnourished child as present in this case, allows clinicians to consider tuberculosis as potential diagnosis worth investigating and managing. The child had a BCG scar, indicative of immunization but this does not sufficiently rule out tuberculosis of the vertebra. The risk of multiple infections was high since the child was not on routine Penicillin V and had no protection from *Hib* and *Streptococcus pneumoniae* vaccines. He was therefore at risk of overwhelming...
encapsulated infections [23,24]. The lack of blood culture results was a major limitation in addressing the role bacterial infection contributed to the development of the pathologies. Nevertheless, patient responded to a course of antibiotics and other treatments provided.

Aseptic necrosis of the large bone heads is a common manifestation in the older SCD patients [25,26]. The progression of aseptic necrosis is facilitated by malnutrition and secondary bacterial infections. This child presented with significant level of malnutrition (Weight for Height z-score < -3SD), osteopenia and evidence of joint space narrowing are clearly risk factors for progressive aseptic necrosis. Similar explanation above could account for the pathological rib fractures and the gibbus observed in this child. Acute Chest Syndrome (ACS) is one of the life threatening conditions of SCD and is usually confused with pneumonia in children [27,28]. Clinicians should have high index of suspicion for ACS. This child did not present with ACS but radiological evidence of opacities in the lung field demonstrates past occurrence of ACS. DVT is not uncommon disease in children and cases have been reported [29,30]. The occurrence of this disease is associated with risk factors such as cancers, trauma, infections, venous catheterization, sickle cell disease and coagulation disorders [29]. This patient had two risk factors for DVT; Sickle Cell Disease and possible multiple infections [31]. In addition to these risk factors, was the fact that this child was immobile for most of the time during the onset of the illness (suffering from septic arthritis of the hip and pyomyositis). In SCD patients, presentation with lower limb pain and tenderness is mostly associated with vaso-occlusive episodes and/or osteomyelitis in areas with high prevalence of the disease. Clinicians should have a high index of suspicion for DVT since it can occur in cases with multiple risk factors. This is critical since the occurrence of Pulmonary Embolism (PE) from the DVT will be life threatening and could also be easily confused to be ACS in a SCD patient and patient would be mismanaged. This short case report therefore tried to highlight some of the uncommon pathologies amongst others that might arise in a SCD patient and to alert clinicians to look out for DVT as an uncommon differential diagnosis in swollen limbs in these patients. It is worth noting that attention needs to be paid to a multi-disciplinary preparedness and response approach to the management of SCD patients, as presented in this case report, to mitigate or minimize disability and mortality.

References


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