Adrenal Myelolipoma, Cholelithiasis and Calcified Spleen: Retrospective Diagnosis of Sickle Cell Anemia Using a Novel Triad of Abdominal Imaging Findings

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ABSTRACT
Sickle cell anemia is an inherited abnormality of the globin chain with very high prevalence in the Indian subcontinent. A significant proportion of these patients present late in life and are at a risk of complications like acute chest syndrome and painful episodes till a definitive diagnosis is reached and appropriate treatment is started. We report a novel triad of abdominal imaging findings which is not reported in literature until now and which may suggest a diagnosis of sickle cell anemia in retrospect. Patients with this triad of abdominal findings should be suspected to have an underlying hemoglobinopathy and should be referred for further hematological workup. Although in our case the patient was diagnosed to have sickle cell anemia depending on the abnormal morphology of red cells and hemoglobin electrophoresis, it should be remembered that this triad of findings may be seen in other hemoglobinopathies which induce a state of chronic anemia.

Key words: sickle cell anemia, adrenal myelolipoma, gall stones, calcified spleen.

INTRODUCTION
Adrenal myelolipoma is a rare non functional benign tumor which is incidentally diagnosed in asymptomatic patients in the 4th and 5th decades. It has a prevalence of approximately 0.08-0.4% on autopsy findings. The coexistence of adrenal myelolipomas and cholelithiasis is indeed very unusual and has rarely been documented. In a
recent article published in 2011 Dahiya et al. stated that this unusual association has been reported only twice prior to their publication. We found another case report describing this unusual association taking the total count to four prior to our publication. However an extensive search for literature did not reveal any reports of association of a shrunken calcified spleen along with adrenal myelolipoma and gall stones and this is probably the first case report of this unusual triad of findings. We present a case of a 28 year old female who was retrospectively diagnosed to have sickle cell anemia based on this triad of findings on abdominal imaging.

CASE ILLUSTRATION

A 28 year old female presented with right hypochondriac pain for the past 6 months which was not associated with retrosternal burning pain or vomiting. On palpation there was mild tenderness in the right hypochondriac region and mild pallor was seen in the conjunctiva, otherwise, rest of the systemic examination was within normal limits. An abdominal ultrasound study revealed multiple gallstones ranging in size from 0.5 cm-1 cm. The gall bladder wall appeared normal and there was no evidence of any peri cholecystic fluid collection. A fairly large soft tissue mass was seen in the right supra renal region measuring approximately 6.9 cm X 5.5 cm in maximum dimensions. The mass was seen to be separate from the liver and the right kidney, however, the right adrenal gland could not be visualized separately. The mass appeared predominantly hyperechoic in echo texture with few hypoechoic areas within it. There was no calcification within the mass and Doppler evaluation did not reveal any significant vascularity within the mass. A neoplastic mass arising from the right adrenal gland was suspected and the patient was referred for CT scan of the abdomen. CT scan showed a soft tissue mass in the right supra renal region measuring approximately 7 cm x 5.8 cm [Figure 1, 4 (short arrow)]. The right adrenal gland could not be identified separately. The mass showed ill defined areas of soft tissue attenuation (-5 HU) were also seen within the mass. Based on these characteristic imaging findings, working diagnosis of right adrenal myelolipoma was made. In addition multiple hyper dense calculi were seen in the gallbladder without any radiological evidence of cholecystitis [Figure 2]. The spleen could not be visualized in the splenic fossa. A densely calcified mass was seen in the splenic fossa which was presumed to be a chronic infarcted calcified spleen [Figure 3, 4 (long arrow)].
Blood examination revealed hemoglobin of 8.4 gm/dl and red blood cell count of 3.02 x 10^6/microlitre. The mean corpuscular volume, MCH and MCHC were 81.7 fl, 27.8 pg and 34.0 gm/dl respectively. Few target cells and sickle cells were also seen. Hemoglobin electrophoresis revealed an Hb S of 86.4%, Hb A2 of 3.3% and Hb F of 6.5% thereby confirming a diagnosis of sickle cell anemia. At the time of submitting the case report for publication the patient is undergoing pre surgical workup for removal of the adrenal tumour.

DISCUSSION

Sickle cell anemia is an inherited abnormality of the beta globin chain due to substitution of valine for glutamic acid resulting in abnormal shaped (sickled) red blood cells (RBC). These deformed RBC’s have increased propensity for microvascular occlusion leading to ischemia and infarction in various body organs. As the RBC’s have abnormal shape, they are removed earlier from the blood circulation resulting in hemolytic anemia.

The highest frequency of sickle cell disease is found in the tropical regions; particularly sub Saharan Africa, India and Middle East. The prevalence of sickle cell disease in India has ranged from 9.4% to 22.2% in endemic areas with India alone accounting for 50% of cases all over the world. The association between adrenal myelolipoma and haemoglobinopathies including thalassemia, sickle cell anemia and hereditary spherocytosis has been proposed earlier by a few authors although there might definitely be more complex pathogenetic mechanisms underlying this entity.

Adrenal myelo lipomas are rare benign tumors composed of adipose tissue and myeloid elements. The association of myelolipoma with obesity, hypertension and malignancies has been described earlier however an association with gallstones has been described in only 4 cases prior to our publication. A chronically infarcted calcified spleen along with adrenal myelolipoma and gallstones on computed tomography has not been reported till now to the best of our knowledge. Dahiya et al stated that the co occurrence of gall stones and adrenal myelolipomas may be an incidental finding or there may be a pathogenetic mechanism which may be yet unknown. Pleuripotent hematological stem cells are present in the embryonic mesonephros which subsequently differentiates to form the adrenal cortex. Metaplasia of these embryonic nests of stem cells may give rise to the myeloid elements within the adrenal. Occurrence of myelolipomas in association with hemoglobinopathies suggests that the myeloid elements in adrenal gland are under the influence of erythropoietin which is produced in abundance due to the chronic anemic state. Thus extra medullary hematopoiesis within the adrenal may contribute to the development of myelolipomas.

Approximately 50% of adults and 20% children with sickle cell anemia patients have gallstones and 94% of patients undergo autosplenectomy by the age of 5 years. However the exact incidence of adrenal myelolipomas in patients with sickle cell anemia is not known with only a few case reports documented in literature. Most probably the stimulation of adrenal myeloid cells is a chronic process which occurs over a long period of time as in our patient who was not diagnosed with sickle cell anemia up to the age of 28 years. The late presentation and chronic anemia also explain the calcified spleen that must have undergone repeated infarctions over a long period of time ultimately resulting in a calcified avascular mass.

It is important to recognize this triad of findings on abdominal imaging and refer these patients for appropriate hematological workup. Approximately 61% of patients with sickle cell...
anemia present by the end of the first year, 78% by the end of second year and 86% by the end of the third year. The remaining 14% present by the age of 10 although late presentation by the age of 20 years is also known. A sizeable number of patients present late in life and are at risk of complications until a definitive diagnosis is achieved and treatment is initiated. The mainstay of treatment of sickle cell anemia is hydroxyurea, an oral chemotherapeutic agent that acts by reducing the production of red cells containing a high concentration of sickle hemoglobin and favoring the production of red cells containing high fetal hemoglobin levels. A Multicenter Study of Hydroxyurea in Sickle Cell Anemia conclusively showed that, over 2.5 years, hydroxyurea diminished the morbidity of SCA in adults with frequent painful episodes by reducing the incidence of painful episodes and acute chest syndrome by nearly half.

**CONCLUSION**

Young adults with sickle cell anemia are a vulnerable population with a high risk of mortality at the interface between pediatric and adult medical care. Quinn et al even argued for creating special clinics for adolescent and young adults with SCA, run jointly by pediatric and adult medical providers because young adults seem to be at high risk of death. It is of utmost importance that the diagnosis of SCA be made at the earliest and this novel triad of abdominal findings should alert the radiologist to refer the patient for further detailed hematological workup. Although in our case the patient was diagnosed to have sickle cell anemia depending on the abnormal morphology of red cells and hemoglobin electrophoresis, it should be remembered that this triad of findings may be seen in other hemoglobinopathies also which induce a state of chronic anemia.

**REFERENCES**