Dilated Cardiopathy Associated with Sickle Cell Disease in a 68 Years Old Female: An Emerging Complication in Sub-Sahara Africa

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To cite this article:
Sylvie Ndongo Amougou, Mary Anne Ngam, Murielle Helles Lema, Mazou Ngou Temgoua, Aicha Yap Mefire, Anderson Ngouo Tchiffo, Samuel Kingue. Dilated Cardiopathy Associated with Sickle Cell Disease in a 68 Years Old Female: An Emerging Complication in Sub-Sahara Africa. Cardiology and Cardiovascular Research. Vol. 3, No. 3, 2019, pp. 65-70. doi: 10.11648/j.ccr.20190303.15

Received: August 23, 2019; Accepted: September 9, 2019; Published: September 21, 2019

Abstract: Sickle Cell Anemia (SCA) is an autosomal recessive disease caused by a point mutation in the hemoglobin beta gene found on chromosome 11p15.5 [1]. Specifically, it occurs when a single base from A to T in the codon for glutamic acid at position 6 is changed to valine of the beta globin and thus disrupts the tertiary structure and stability of the hemoglobin molecule [2]. Sickle hemoglobin is responsible for wide spectrum of disorders which vary with respect to severity of anemia, frequency of crises and duration of survival [3]. We present the case of a dilated cardiopathy in an elderly female sickle cell patient. A 68 years old female sickle cell patient with no known major cardiovascular risk factor presented with progressive onset of dyspnea. Clinical examination showed signs of left ventricular failure; an electrocardiogram showed a sinus regular rhythm, left ventricular hypertrophy with systolic overload, Q waves in the anteroseptal leads. Cardiac ultrasound showed a dilated cardiopathy with a preserved left ventricular systolic function at 57%, normal regional wall motion and normal pulmonary pressure. We concluded of heart failure due to probable chronic anemia or anischemic cardiopathy with a conserved systolic function in an elderly sickle cell patient but we were not able to confirm the main etiology without CT Coronary Angiogram or coronarography. Sickle cell anemia is a common genetic condition in sub-Saharan Africa associated with early death. This case is special because we have an elderly female presenting with heart failure on a dilated cardiopathy. This enhances the necessity of strict cardiovascular follow up of Sickle cell patient.

Keywords: Sickle Cell Disease, Dilated Cardiopathy, Older Female, Sub-Sahara Africa

1. Background

Sickle Cell Disease (SCD) is a blood disease, which is caused by an inherited Hemoglobin S (HbS) gene. It is caused by a point mutation in the 6th codon of the beta globin gene leading to the substitution of glutamic acid by valine [4]. The genetic mutation described causes polymerization of the hemoglobin molecule that alters the erythrocyte shape and its ability to deform. There is an increased adhesion of erythrocytes followed by formation of hetero cellular aggregates, which physically cause small vessel occlusion and resultant local hypoxia. This process triggers a vicious cycle of increased HbS formation, the release of inflammatory mediators and free radicals that contribute to reperfusion injury. Hemoglobin also binds to nitric oxide (NO), a potent vasodilator, and releases oxygen. Erythrocytes are more likely to sickle and become rigid in the presence of dehydration.
Other associated pathological events include increased neutrophil adhesiveness, nitric oxide binding, increased platelet activation, and hypercoagulability. [5]. Sickle cell anemia is a multi-system disorder. Patients are completely asymptomatic before the age of 6 months due to the presence of fetal hemoglobin which gradually decreases and HbS becomes predominant [6]. The most common presenting feature is vaso-occlusive crises. Patients commonly complain of excruciating pain in the abdomen, thorax, joints, long bones, and digits. Some individuals may experience multiple episodes while others may remain free of them for long periods of time. Signs and symptoms of anemia are also prevalent, including palpitations, fatigue, pallor, and tachycardia. Repeated vaso-occlusive crises may result in splenic infarctions and resultant functional asplenia. This asplenia results in repeated infections with encapsulated bacteria like *Streptococcus pneumoniae*, *Staphylococcus aureus*, and many others. These pathogens may cause life-threatening pneumonia and septicemia which are usually fatal. Aplastic crises are another significant manifestation of sickle cell disease. Here, the presence of parvovirus B19 challenges an already stressed bone marrow, it fails to generate the appropriate number of RBCs which results in severe anemia. It usually last for 5 to 7 days but can also be life threatening [5-6]. The introduction of treatments that induce protective fetal hemoglobin and reduce infectious complications has greatly prolonged survival. However, with increased longevity, cardiovascular complications are increasingly evident, with the notable development of a progressive proliferative systemic vasculopathy, pulmonary hypertension (PH) and left ventricular diastolic dysfunction. Also, chronic anemia results in cardiac chamber dilation and a compensatory increase in left ventricular mass. This is often accompanied by left ventricular diastolic dysfunction [7]. We present the case of a dilated cardiopathy in an elderly female sickle cell patient.

2. Case Presentation

A 68 years old woman who is a retired secretary, married with four children and known homozygous sickle cell patient since infancy. Her hemoglobin level is usually between 6 to 7 g/dl. She has a history of repeated blood transfusion and folic acid supplementation due to severe anemia. She had vaso-occlusive crises monthly before the age of 40 years and after 40 years she had crises at least once every 3 years. She is on folic acid 5mg/day and deferoxamine, an iron-chelating agent. She was also diagnosed of hepatitis C infection with a viral load of 7021095 U/L and successfully treated at the age of 40 with ledipasvir and sofosbuvir. She presented with a 2 weeks history of progressive onset of dyspnea first on exertion and later evolving to dyspnea at rest associated with palpitation and intense asthema. There were no associated joint pains, fever or loss of consciousness. Physical examination showed an ill looking conscious and oriented patient. Patient had signs of respiratory distress that is oxygen saturation of 89%, tachypneic at 40 cycles per minute. Also, she was, tachycardic at 105 beats per minute apyrexic at 37.8°C with a blood pressure of 125/85mmHg. There were signs of left ventricular failure that is gallop rhythm, bilateral basithoracic lung crepitation and those of right ventricular failure such as distended jugular vein, hepatojugular reflex and a right ventricular heave. There was no tenderness or redness over any joint surface. A diagnosis of a probable decompensated global heart failure was made.

Paraclinical workup done showed

Resting electrocardiogram Sinus regular rythm, left ventricular hypertrophy with systolic overload, significant Q waves in the antero-septal region (V1, V2, V3), (Figure 1).

![Figure 1. Resting electrocardiogram.](image-url)
Transthoracic Doppler ultrasounds (Figures 2, 3, 4, 5) showed dilated left ventricle, eccentric left ventricular hypertrophy with a conserved left ventricular systolic function at 57%, grade II diastolic dysfunction, normal pulmonary pressure with PAPS at 24mmHg and dilated left atrium. There was no thrombus inside the cavities; valves were of normal morphology and function.

Figure 2. Transventricular Parasternal Long axis view.

Figure 3. 4 cavities view showing enlargement of the left atrium.
Figure 4. Trans-aortic Parasternal long axis view showing normal diameter of the Aorta root.

Figure 5. Mitral Profile in apical 4 cavities view.
Biological workup (lipid profile, renal function test cardiac enzymes) were normal. She had an elevated ferritin level (2076; normal range: 9-120 ng/ml) and severe normocytic hypochromic anemia (HB: 6.5 g/dl, Normal range: 12-16 g/dl; MGV: 83, range: 80-96 fl; MCHC: 24, range: 27-32 pg).

Due to limited finances, we were unable to do CT Coronary Angiogram, coronarography, Brain natriuretic peptides (BNP) and Pro brain natriuretic peptides (NT Pro BNP). Dobutamine stress echocardiography was difficult to do because of the risk of inducing pain. Aworking diagnosis of global decompensated heart failure NYHA IV on a dilated cardiopathy of multifactorial origin (ischemia, anemia, hyperferitinemia, HVC or toxic due to past-exposition to anti-HVC drugs) with conserved of Ejection fraction at 57% was made. Patient was admitted in an intensive care unit where patient received as treatment Oxygen starting at a rate of 4L/min with gradual increase of rate based on the Oxygen saturation; Fluid restriction at 1 L/min; Loop diuretic (Furosemide at a dose of 1mg/Kg); Angiotensin converting enzyme inhibitor (Ramipril 1.25mg per day). Evolution on this treatment was marked by regression of signs of congestion (patient became less dyspneic with an improvement of oxygen saturation). The dose of the diuretic was reduced in other to prevent dehydration which could lead to a vaso-occlusive) and low dose aspirin (100mg per day) with beta blocker (Metoprolol 50mg per day) were added to the treatment.

3. Discussion

Sickle cell disease (SCD) is a common genetic disorder with potentially devastating consequences for those affected [8]. It is caused by a point mutation in the 6th codon of the beta globin gene leading to the substitution of glutamic acid by valine [4]. In 1994 at boston, Platt and al. found the median age of death was 42 years for males and 48 years for female [12]. Sophie Lanzkron and al. used data of the National Center for health Statistics of U.S. to find that the period being under 5-years old [9]. Our patient is an elderly female patient who has survived up to her 60s despite severe vaso occlusive crises and severe anemia. Also, she was able to have four successful pregnancies with normal term deliveries. Her past history of Hepatitis C could be due to the repeated blood transfusion as blood transfusion is a risk factor of hepatitis C infection [10]. Viral hepatitis C is a systemic disease which can have extra hepatic manifestations either due to the virus or the treatment. Amongst them we have cardiovascular manifestations such as ischemic cardiopathy [15-17]. But she was treated since more than 10 years; we have not found studies which show for how long we can have cardiovascular effects after the treatment.

Patients with sickle cell anemia usually have multiple organ involvement. In SCD, a number of the chronic complications appear to be related to hemolytic anemia while other complications are related to inflammation and vaso-occlusion (classic “sickling” events) [11]. Vascular occlusion of small and large vessels can lead to chronic damage of multiple organs including brain, lung, bone, kidney, liver, spleen, and retina. However, the extent to which SCD impacts myocardial function is not very clear. Cardiovascular manifestations include cardiac chamber dilatation because of chronic anemia with the increase in cardiac output, both right and left ventricular systolic and diastolic dysfunction, elevated cardiac output, pulmonary hypertension, cardiomegaly and myocardial ischemia. Progressive heart damage from iron overload occurs in patients requiring routine transfusion therapy. Pulmonary hypertension resulting from intravascular hemolysis has also been recognized as a major complication that independently correlates with survival [7, 18]. The echocardiography of the patient showed dilated left ventricle and atrium, diastolic dysfunction but normal right ventricle and pulmonary pressure. The normal pulmonary pressure, a good observance of the treatment, a knowing of the vaso-occlusive crisis may be some of the reason why the patient lives more than the others.

Ferritin level was elevated probably due to multiple transfusion and iron supplementation. Grossly visible cardiac iron deposits are associated with cardiac dysfunction and usually with chronic cardiac failure. Diastolic dysfunction appears early in the course of iron overload while systolic dysfunction occurs very late. Iron accumulation occurs initially in the ventricular myocardium followed by the atrial myocardium, but remains greater in working than in conducting myocardium. The degree of the heart block (first degree) and supra-ventricular arrhythmias are correlated with the level of iron deposition in the atrial myocardium [18]. In the past, our patient had history of multiple arrhythmias with bigeminism in the electrocardiography.

In this patient, the ischemic cardiopathy could be due to an obstruction of a coronary artery by sickled hemoglobin since the patient did not have major atherosclerotic risk factors. The patient did not present with current signs and symptoms of an acute coronary syndrome because the obstruction of the coronary artery might have occurred in the past and might have being treated as an acute chest syndrome. The treatment instituted in this patient was meticulous since as the patient had to be depleted with provoking dehydration which could lead to a painful vaso-occlusive crises. Our limitation was the inability to do a coronary angiography in other to exclude the presence of coronary artery plaques.

This case shows the longevity of a female homozygous sickle cell patient who had a normal life style presenting in
heart failure due to multifactorial cause. The treatment was meticulous such that the patient was depleted without provoking dehydration which could lead to painful vaso-occlusive crises.

4. Conclusion

Heart failure associated to SCD was previously considered as rare because of reduced life expectancy. With advances in modern treatment, long term complication become frequent and the clinician should pay attention of that in follow up of the patient. This case enhances the necessity of strict cardiovascular follow up of Sickle cell patient.

Abbreviations

BNP: Brain Natriuretic Peptide
HbS: Hemoglobin S
MCHC: Mean Corpuscular Hemoglobin Concentration
MGV: Mean globular volum
NYHA: New York Health Association
NO: Nitric Oxide
PH: Pulmonary hypertension
RBC: Red Blood Cell
SCA: Sickle Cell Anemia
SCD: Sickle Cell Disease
US: United States

Authors’ Contributions

NSA managed the patient, ANT and AYM participate in the echographic procedure. NME, HML drafted the initial manuscript which was modified by MNT. SK supervised all the process. All authors read and approved the final manuscript.

Consent for Publication

Written informed consent was obtained from the patient for publication of this case report and any accompanying images. A copy of the written consent is available for review by the Editor-in-chief of this journal.

Competing Interests

The authors declare that they have no competing interests.

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