Pulmonary artery aneurysm in Behcet’s disease: a ticking time bomb

Abstract

Behcet’s disease (BD) is a rare condition in sub-Saharan Africa. It often manifests as recurrent oral and genital aphthous ulcers with ocular features and infrequently involving major systems like pulmonary vasculature and parenchyma. Pulmonary involvement in BD, although rare, can manifest as pulmonary artery aneurysm, vascular thrombosis, pulmonary infarction, pneumonia and pleurisy.

Our index case is a 38 year old Nigerian who presented with recurrent orogenital ulcer, hemoptysis and uveitis. He had a contrast enhanced chest CT scan that revealed pulmonary artery aneurysm. His symptoms resolved following commencement of immunosuppressant and steroid.

Increased awareness is required for early detection of BD especially in regions where it is regarded as rare. Furthermore, multidisciplinary approach in its management is vital to reduce morbidity and mortality.

Keywords: behcet’s disease, pulmonary vasculitis, pulmonary artery aneurysm, Nigeria

Introduction

Behcet’s disease is a rare multi systemic disorder of unknown etiology which often manifest with recurrent oral aphthous ulcers, genital ulcers and ocular lesions. It occasionally manifest as severe systemic presentations involving the pulmonary, cardiovascular system, central nervous system, and gastrointestinal system.¹

Behcet’s disease (BD) has a worldwide prevalence which ranges between 0.1/1000 and 1/10,000 with a significant presence in Asian countries from the Mediterranean to Japan, hence the term Silk Road Disease.² Turkey has reported the highest prevalence ranging from 80 to 420/100,000. However, there are few reported cases in Africa, North Europe and America.³,⁴ Males in western Asia are commonly affected as opposed to females in Japan and Korea. The usual age at presentation is between the third and fourth decades.²

Although the underlying cause of BD is unknown, like other autoimmune disorders, it is postulated to occur in genetically predisposed individuals exposed to triggering agents like microbes and environmental factors. The presence of an HLA-B51 allele is associated with increased risk of developing this condition as well as worsened presentation.⁵,⁶

The clinical presentation is highly variable and no specific laboratory investigation is available for diagnosing BD. Several diagnostic criteria were proposed in the last 70years with different racial considerations but none is specific to individuals of African descent. However, two separate consensus criteria made by the collaboration of experts from several countries; International Study Group on Behcet’s Disease (ISG) and International Criteria of Behcets Disease (ICBD), have received wide acceptance with varying sensitivity and specificity.⁷ The use of anti-inflammatory and immunosuppressive agents is important in relieving symptoms as no curative solution is currently available.⁸

Case presentation

A 38 year old Nigerian male who presented to the respiratory clinic with a history of chronic cough productive of blood stained sputum of 12months duration, recurrent oral and genital ulcers of 12months, low grade fever of 10months and 8months history of progressive weight loss. Mouth ulcers were recurrent and painful, involving the tongue and gum. The genital ulcers were multiple and painful with no bleeding or discharges and healed spontaneously without scars. He had had about 5 recurrences in the last 12months. There was associated history of blurring of vision and occasional redness of the eyes with no eye discharge. The patient reported no history of joint pains or swelling. He denied history of contact with adult with chronic cough or past treatment for pulmonary tuberculosis or any history suggestive of sexually transmitted infections in the past. He also denied use of sculpture containing drugs and occupational or environmental exposure to toxic agents. His other family members were healthy and none had history of similar illness. He recently got married but his wife is yet to conceive.

Physical examination revealed a wasted young man (BMI 18.1kg/ m²) who was afebrile with no lymphadenopathy. He had multiple tender oral ulcers measuring about 1-2mm in diameter on the tongue and gum with an erythematous base. However, normal male genitalia with healed without scars and there was no oedema or tenderness of the testicles. Ophthalmologic examination revealed normal visual acuity and optic nerve with features of uveitis on slit lamp examination.

The results of laboratory investigation revealed packed cell volume (38%). Other cells counts were within normal limit but for the peripheral blood film that showed microcytes, macrocytes and anisocytosis. Chest radiograph showed widespread rounded opacities in both lung fields while constrast enhanced CT scan revealed bilateral multiple saccular dilatations of the pulmonary arterials in keeping. Sputum for acid and alcohol fast bacilli and Gene expert (negative), erythrocyte sedimentation rate (40mm/hr), Hepatitis screening (HBsAg, Anti-HCV were both negative), HIV rapid serology test (negative), and VDRL for Syphilis (negative). The values of Anti-Nuclear Antibody, pANCA, cANCA were within normal limit and pathergy test was negative after 48hrs.
An assessment of Behcet’s disease was made based on the presence of recurrent oral ulcer (>3 times in a year), recurrent genital ulcers and posterior uveitis as proposed by International Study Group on Behcet Disease. He was commenced on oral prednisolone 10mg bid, oral azathioprine 50mg bid and omeprazole 20mg b.d. The dose of the oral prednisolone was tapered off after 2 weeks. He made significant clinical improvement as all symptoms resolved after 2 weeks of treatment and complete resolution of symptoms was noticed after a month (Figures 1-3).

Figure 1 Black arrows indicate areas of multiple ulcers on the tongue.

Figure 2 Chest X-ray: arrows show rounded opacities seen in both lung fields.

Figure 3 Shows a contrast enhanced Chest CT of patient that revealed bilateral multiple saccular dilatations of the pulmonary arterial branches.

Discussion

Behcet’s Disease (BD) is a multi-systemic disorder of unknown etiology, rarely reported in sub-Saharan Africa, characterized by recurrent attacks of oral aphthous ulcers, genital sores, and ocular manifestations. It runs a severe course in men and those with onset before 25 years of age. Vascular involvement is the most common cause of mortality and associated with poor prognosis. Its pulmonary manifestations could mimic conditions like pulmonary tuberculosis, especially in tuberculosis high burden countries.

Our index patient is a male in his fourth decade with recurrent oro-genital ulcers and uveitis. This is in tandem with the diagnostic criteria for BD as well as the archetypal chest CT finding of pulmonary artery aneurysm. The age of onset in our patient is higher than the 25 years associated with poor prognosis, however, the gender as well as the evidence of pulmonary artery aneurysm has been associated with high mortality and morbidity.

Pathergy test is the hyper-reactivity of the skin to intracutaneous injection or needle prick but this was negative in our patient. It is demonstrated more frequently in specific population with rare occurrence in African studies. This could be responsible for the negative pathergy test in our patient as HLA B51 influences clinical features of BD in susceptible individuals.

ANCA-associated vasculitis that mostly affect the lungs were excluded as connective tissue disease serologies and anti-neutrophil cytoplasmic antibody (ANCA) were normal. Similarly, Hughes-Stovin syndrome (HSS), a close differential of BD with consistent radiologic and histopathologic finding was unlikely as our patient had both oral and genital ulcer which are usually absent in HSS. Patient could have been managed as smear negative tuberculosis in a high burden country but for the rapid and sensitive diagnostic test of Xpert MTB/RIF test which was also negative.

Our patient was commenced on immunosuppressive agents and steroid as advocated by the European League against Rheumatic Diseases (EULAR) and there was significant clinical improvement within 2 weeks with no side effects. Recommended treatment for pulmonary aneurysm is mainly with immunosuppressives as surgery is associated with high risk of mortality. Furthermore, patients with ocular involvement benefit from azathioprine and steroid to prevent irreversible damage and visual loss.

Conclusion

It is apparent that the management of Behcet’s disease may be challenging to the attending physicians, especially among populations with documented rare occurrence. There is, therefore, a need to increase the level of awareness about BD among physicians for prompt diagnosis and treatment of individuals with BD so as to prevent disability and reduce associated mortality.

Consent

All authors declare that written informed consent was obtained from the patient.

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None.
Conflict of interests
The author declares no conflict of interests.

References