Klippel-Trenaunay Syndrome: A Case Report and Literature Review

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Abstract: Klippel Trenaunay Syndrome (KTS) is a rare, sporadic vascular malformation characterised by its clinical triad of: port wine stains, varicosities and limb hypertrophy. Here, we present a case of a 51-year-old woman who presented with these symptoms. We will discuss the hypothesized etiology and recommended management of KTS with a literature review.

Keywords: Klippel-Trenaunay Syndrome

1. INTRODUCTION

The eponymously named Klippel-Trenaunay Syndrome (KTS) is a rare cutaneous vascular condition that was first described in 1900 by two French physicians Klippel and Trenaunay, who first noticed this syndrome in two patients who presented with a port wine stain and varicosities of an extremity associated with hypertrophy of soft and bony tissue. They termed the syndrome “naevus vasculosus osteohypertrophicus”. In 1907, Frederick Parkes Weber, unaware of Klippel and Trenaunay’s report, described a patient with the three aforementioned symptoms as well as an arteriovenous malformation of the affected extremity. He termed the process hemangiectatic hypertrophy and more commonly known today as Parkes Weber Syndrome (PWS). KTS must be distinguished from PWS which has a hallmark feature of AVFs [1].

2. CASE REPORT

A 51 year-old Caucasian lady presented to the Vascular Surgery clinic for left lower limb varicosities. She complains of extensive skin stains since birth, with worsening left lower limb swelling and varicosities for the past 20 years. She denies any prior superficial thrombophlebitis, deep vein thrombosis, pulmonary embolism, venous ulcerations or rectal bleeding. As she is a home-maker, the limb swelling and varicosities do not affect her daily activities. Upon examination, the patient has extensive port wine stains over her left arm, torso and left lower limb which were associated with left lower limb varicosities and left thigh hypertrophy (Figures 1 to 4). The patient has the Klippel-Trenaunay Syndrome clinical trial of port wine stains, varicosities and limb hypertrophy. Ultrasound did not reveal deep vein thrombosis and as patient was not keen for further surgical intervention, she did not proceed with further venous insufficiency Ultrasound Duplex.

Figure 1. Left upper limb portwine stains

Figure 2. Left lower limb port wine stains, hypertrophy and varicosities
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3. DISCUSSION

KTS affects men and women equally and is not limited to any racial group. Several molecular characteristics have been identified- including a translocation of chromosome 8q22.3 and 14q13 [2], which is associated with vascular and tissue overgrowth syndrome while others have found a terminal deletion of 2q37.3 in a patient with KTS [3]. Another study identified found an association of the vascular and/or overgrowth gene location on chromosome 18 [4].

KTS is typically characterized by a clinical triad of extremity varicosities, cutaneous vascular malformations, and hypertrophy of soft tissues and long bones. In a study of 252 patients with KTS (136 females and 116 males), capillary malformations in the form of port wine stains were found in 98% of patients (246 patients), 72% of the patients had varicosities (182 patients), and an incidence of soft tissue hypertrophy in 67% of the patients (170 patients). From the results, the student concluded that only 63% (159 patients) of the patients studied had the classic clinical triad while a much fewer number of 93 patients (or 37%) had two out of the three features of KTS [5].

Most KTS patients present with unilateral lower limb involvement (85%) while 12.5% present with bilateral lower limb involvement and rarely, crossed bilateral involvement in 2.5% of the patients [6]. Hence, some has proposed the method of using a venous ultrasound duplex study as well as nuclear and radiological imaging to further confirm the diagnosis, as the clinical triad may not be present in all patients [7]. A venous ultrasound duplex study will not only help to confirm the presence and determine the extent of affected vessels, but also assess for complications of venous thrombosis. To differentiate between KTS and PWS, segmental limb pressure and ankle brachial index studies can be done to rule out the presence of arteriovenous shunts which is characteristic of PWS and not present in KTS [5]. With age, symptoms of varicosities and pain may worsen and can progress to the most severe complication of venous thromboembolism and pulmonary embolism [8].

Additional imaging to assess complications of KTS such as plain X-ray for imaging of long bones in patients with limb length discrepancy. Another useful tool for non-radiological or non-contrast imaging which would allow for better imaging of the different tissues would be a magnetic resonance (MR) scan or a MR angiography. It has the additional benefit of identifying lymphedema.

A multidisciplinary approach as well a nonsurgical approach has been the mainstay of KTS management. Treatment is largely symptomatic control and complication prevention. Complications such as venous thromboembolism and cellulitis through anticoagulation and the use of compression stockings [9]. Surgery is rarely indicated unless only in symptomatic patients who may require epiphysseodesis, ligation or stripping of incompetent veins. Additional surgical technique also includes subfascial endoscopic perforator surgery (SEPS).

4. CONCLUSION

Klippel Trenaunay Syndrome (KTS) is a rare, sporadic vascular malformation characterised by its clinical triad of: port wine stains, varicosities and limb hypertrophy. A multidisciplinary approach is the mainstay of KTS management and treatment is largely symptomatic control and complication prevention.

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