Transient thromboembolic event in a child with severe hemophilia A due to venous obstruction

Athugalpura DMAA, Jayathilaka MM, Jayathilaka A, Gunathilaka PKG, Kankananarachchi I, Gamhewage N

Abstract

Venous thrombosis is a very rare occurrence in patients with hemophilia A. We present a one year old child with severe hemophilia A, who developed deep vein thrombosis in antecubital fossa following venous obstruction due to bleeding around the vein. We didn’t start anticoagulant therapy to our patient because of the risk of further bleeding into subdural hemorrhage he had already developed. We could manage the thrombosis and save the limb by relieving the obstruction with replacement of factor VIII.

Introduction

Hemophilia A is a hereditary hemorrhagic disease characterized by deficiency of coagulation Factor VIII. Venous thrombosis in patients with hemophilia A has been rarely encountered in medical literature [1, 2, 4-7] and it is an important cause of hospital acquired morbidity and mortality [3]. In this case report we present the onset of deep vein thrombosis in a one year old child with hemophilia A.

Case report

One year old boy, who had the diagnosis of severe hemophilia A (Factor level 0.286%), and a past history of spontaneous intra cranial hemorrhage at 8 months of age, has presented with 2 days history of fever, reduced activity and drowsiness. CT scan revealed acute sub dural hemorrhage in falx & in tentorium and chronic hemorrhage in left sided parieto-occipital region. He was on prophylactic factor viii weekly prior to current admission because of the previous spontaneous intra cranial hemorrhage. According to guidelines we started giving 100% factor viii correction twice daily to overcome further bleeding into sub dural space.

In spite of these measures, APTT remained elevated between 80 s and 90 s and the child had been less active and irritable. On day 9 of the ward stay, child has developed gross left upper limb swelling and we suspected him as having a local bleed followed by venous obstruction. Doppler ultrasonography showed ‘Deep vein thrombosis in antecubital fossa without evidence of arterial compromise’. The vein at antecubital fossa in which the thrombosis developed was almost completely occluded due to hematoma around it. Activated partial thromboplastin time, bleeding time and other investigations are given in table 1.

We decided to continue factor viii correction without anticoagulation because of the risk of further bleeding. Instead, we increased the dose of factor VIII. In addition, we gave a dose of prothrombin complex concentrate (PCC). At the same time blood was taken for ‘Bethesda test’ because we suspected inhibitors to factor VIII.

Discussion

Venous thrombosis is an important cause of hospital acquired morbidity and mortality [3]. It is associated rarely with patients of hemophilia A [1, 2, 4-7]. Several risk factors have been described in medical literature. There were hardly any cases with normal coagulation test results at the time of thrombosis. The most important risk factor for patients with hemophilia A was taking Factor VIII inhibitor bypassing agent (FEIBA) or recombinant
activated factor VII (rFVIIa) for inhibitors [3]. Other risk factors were congenital prothrombotic conditions, deficiency of protein C and Factor V Leiden [8], following total hip and knee arthroplasty [9], following insertion of central venous catheters [10]. Murat Bicer et al reported a spontaneous deep vein thrombosis originated from the right main and external iliac veins in a 32 year old white male hemophilia A [11]. Van der Planken reported a deep venous thrombosis development within 18 days after recombinant activated factor VII (rFVIIa) infusion in a 38 years old patient with hemophilia A [12]. Ettingshausen et al. described portal venous thrombosis in a patient with the diagnosis of hemophilia A and Factor V G1691A mutation [6].

There are no clear consensuses regarding the management of thromboembolism in patients with hemophilia A because of the low number of cases in medical literature [11]. Dargaud et al. have used unfractioned heparin for a month subsequent to Factor VIII replacement [1]. Low molecular weight heparin was administered for 9 weeks by Kashyap et al [8]; Ettingshausen et al had practiced different way of approach where he applied low molecular weight heparin together with Factor VIII replacement, after unfractioned heparin for 14 days.

Murat Bicer et al initiated oral anticoagulant drug after 48 hours of thrombosis because coagulation tests were normal at that time and Factor V Leiden was absent. Factor VIII levels were replaced during patient’s follow-up. He suggested in his case report, to start oral anticoagulants with close monitoring for hemorrhage. In our case we were reluctant to start any anticoagulant, because APTT was never normal. It remained in very higher value. On the other hand patient had already developed sub dural hemorrhage which can aggravate with anticoagulation. Our aim was to cure the hematoma around the vessel relieving the obstruction. We expected spontaneous resolution of thrombosis when we remove the risk factor for thrombosis. Finally we were able to save the limb without any anticoagulant.

References


9. Matthew IStein, MD; Justin park, MD; Stephen Raterman, MD, Prevention of VTE Following Total Hip and Knee Arthroplasty in Hemophilia Patients: cme Article, MAY 2011 | Volume 34 • Number 5: 389 – 391

