Inhibitor in hemophilia and its management-A case report

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Inhibitor in hemophilia and its management-A case report

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Abstract

Hemophilia is a genetic disease with a deficiency of factor VIII or IX. People with inhibitors require special care and treatment due to complexity of the disease. Treatment is expensive due to high dose infusion and unavailability of factor in developing countries like India. This case report focuses on role of comprehensive care and follow-up of an adolescent with inhibitors.

Key words: Children, factor VIII deficiency, hemophilia, inhibitors

Introduction

Hemophilia is an inherited X-linked coagulation disorder caused by deficiencies of the clotting factor VIII (FVIII: hemophilia A) or factor IX (FIX: hemophilia B). In India, 18,353 people are diagnosed to have hemophilia as per the recent annual global survey report (WFH, Annual Global Survey 2016, 2017). Hemophilia is classified based on percentage of factor levels as mild, moderate and severe in clinical severity. Spontaneous bleeds are more in moderate and severe hemophiliacs compared to people with mild hemophilia (Phadke, 2011).

Inhibitors are the antibodies produced by the body to fight off the foreign protein present in the treatment product. People with severe hemophilia have got more risk of developing inhibitors than mild and moderate hemophilia. Inhibitors occur among 25-30% of children with severe hemophilia A and 1-6% of individuals with hemophilia B. Regular testing for inhibitors should be done among newly diagnosed people with hemophilia between the 1st and 50th days of treatment. Check for inhibitor at least twice a year after the 50th day of treatment, until they have received 150-200 doses. Continue checking for inhibitor once a year after that. Before any major surgery also inhibitor needs to be checked. When the person is not responding to standard treatment usually inhibitor is suspected (WFH, 2014).

Case report

A 13-year old boy with factor VIII deficiency was admitted with swelling of the left knee joint. He was diagnosed with hemophilia at the age of two years and on-demand factor replacement therapy. Initially he was treated with skin traction, analgesics and tranexamic acid. But the symptoms did not subside with the treatment. Blood investigations were done to find out the reason for not responding factor replacement therapy. APTT value was 85.9 Sec and inhibitor screen was positive and further Bethesda assay showed 2.1 BU as low responding inhibitors. Later he was treated with high dose factor. The family belonged to lower socioeconomic status and they had another child with hemophilia. On discharge child was stable and was advised to come for physiotherapy. The nurse coordinator played a significant role in educating the
family members about prevention of injuries, inhibitor and its management and regular follow-up.

Discussion
Inhibitors make the treatment difficult and complicated mainly due to financial constraints and the availability of factors. Inhibitors are diagnosed by activated partial thromboplastin time (APTT) and confirmatory tests like Bethesda assay, or Nijmegen method (Verbruggen, Van Heerde, & Laros-van Gorkom, 2009). Inhibitor levels are categorized as ‘high titre’ (more than 5 Bethesda Units) or ‘low titre’ (less than 5 Bethesda Units). ‘Low responding’ and ‘High responding’ is one of the classifications of inhibitors based on how the person’s immune system reacts to factor concentrates (Giuffrida, et al., 2008).

Treatment for a person with the inhibitor is decided based on inhibitor titre and anamnestic response. We also need to consider the site and severity of the bleed. Factor concentrates can be administered at higher doses and/or more frequent intervals for acute bleeding in low responders. Acute bleeding in people with high titre inhibitors are treated with bypassing agents, such as activated prothrombin complex concentrates (APCC) and recombinant factor VIIa (Novoseven) (Franchini & Mannucci, 2011). Factor Eight Inhibitor Bypassing Agent (FEIBA®) is an example for APCC which is made from human plasma and contains variable amounts of clotting factors. This is given once in eight to twelve hours but should be limited to low doses such as 100 U/kg per infusion and no more than 200 U/kg/day. Blood clot formation is a risk which has been associated with excess use of FEIBA. High cost and unavailability of the treatment products in every country are the two main difficulties for patients with inhibitors.

rFVIIa (NovoSeven®) is a synthetic product and needs to be administered once in two to three hours. An antifibrinolytic drug such as Tranexamic acid also can be used for better outcomes. A single-centre, mixed retrospective and prospective study showed that Immune tolerance induction therapy was successful in treating severe hemophilia A patients (Haya, et al., 2019). Regular rehabilitation program helps the individuals with inhibitors as they have higher risk for bleeding. Rehabilitation program needs to be tailored based on age, joint score and other co-morbidities (Kubota, et al., 2018).

Conclusion
The inhibitors occur later in life in severe hemophilia A and their development is generally associated with a change in bleeding pattern. A range of therapeutic options to manage the inhibitor is currently available and comprehensive nursing care is needed to manage teenagers with inhibitors. They need to be emphasized on pain management, regular treatment and screening of inhibitors.

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Reference

