Dental considerations on the management of Idiopathic Thrombocytopenic Purpura in children: case report

Considerações odontológicas no atendimento de crianças com Púrpura Trombocitopênica idiopática: relato de caso

Viviane Ferreira ROSSIER¹
Stella Maria Coda Pinto Alves Campos VIEIRA²
Ana Lídia CIAMPONI³
Renata de Oliveira GUARÉ³

INTRODUCTION

Hemopathies are categorized in three large groups, according to the nature of the altered hematologic component. Thus, they are divided in disorders that affect erythrocytes (red blood cells), leucocytes (white blood cells) and hemorrhagic disorders, among which are included platelet and coagulation factors alterations¹.

Purpura is a disease included in the large group of hemorrhagic disorders and is distinguished by cutaneous hemorrhage and blood loss from mucous membranes and internal organs, which is always related to vascular or platelet alterations². Thus thrombocytopenia, which is distinguished by the decrease in the number of blood platelets, is included in this large group¹,³⁻⁹.

Platelets are small granular corpuscles measuring 2 to 4μ in diameter found in the bloodstream. Their role in the coagulation process is the production of a loose aggregate of platelets (temporary hemostatic plug) immediately after a lesion in the blood vessel. This plug is maintained and posteriorly converted into a definite fibrin clot¹⁰.

Idiopathic Thrombocytopenic Purpura (ITP) is considered one of the most common disorders in children²⁻¹¹, and the incidence of asymptomatic illness is approximately 3⁻8:100,000 children/year¹². ITP differential diagnosis must be performed in order to distinguish it from other conditions that may cause the symptoms of thrombocytopenic purpura, such as thrombocytopenia induced by medication (barbiturate, quinine, lengthen therapy with glucocorticoids), hereditary thrombocytopenia.
(Bernard-Soulier Syndrome, Aldrich-Schönlein Syndrome), vitamin C deficiency, viral infections (HIV, infectious mononucleosis), autoimmune disorders (systemic Lupus erythematosus), aplastic anemia, acute leukemia, and non-Hodgkin’s lymphoma.

In laboratorial exams, complete blood count shows severe decrease in platelet count bellow 20,000/mm³ in acute IPT and between 30,000/mm³ up to 100,000/mm³ in chronic IPT. In the coagulogram, bleeding time and coagulum retraction are possible alterations. The number of white cells is normal and anemia may be present in some cases.

Currently, IPT is considered an autoimmune disease and its etiology is related to immunological mechanisms such as antibodies and antiplatelet immunoglobulin in the blood. Genetic studies show the correlation of clinical development of IPT and certain genotypes, suggesting a genetic predisposition and modulation of the illness progression. IPT is the result of the increment of the destruction of antiplatelet antibodies (autoantibodies) through cells of the reticuloendothelial system, especially the spleen. The autoantibodies bind to platelets and are eliminated by the spleen, thus the disease severity reflects the balance of the capacity to produce platelets by the bone marrow mega-karyocyte and platelet destruction by the reticuloendothelial system.

IPT is categorized as acute and chronic; the latter is described based on the presence of thrombocytopenia for more than six months before the first signs and symptoms. Acute IPT prevails in children below 10 years old, affecting both genders. It is usually related to a viral infection history before the development of clinical signs with an interval of 2-21 days from the viral infection and the beginning of IPT. There are reports that associate the illness to Epstein-Barr infection, varicella zoster infection and, particularly, after rubella vaccination.

Chronic IPT affects more teenagers than younger children, especially the female gender. There is a greater probability of patients affected by chronic IPT to develop autoimmune diseases, and 1/3 of these patients present laboratorial and clinical symptoms of collagen vascular disease. Clinical signs of chronic IPT are milder and in some cases they can conceal and hamper the diagnosis.

IPT clinical symptoms include petechiae, ecchymoses, hematomas, epistaxis, hematuria, mucinous cutaneous bleeding and occasionally tissual hemorrhage. In general, splenomegaly doesn’t occur, and spleen is only palpable in 10% of the patients with acute IPT. Complications are rare; however intracranial, subglottic, gastrointestinal and genital-urinary hemorrhage may occur.

Medical treatment for IPT patients adopts different intervening forms, according to individual conditions of each patient given that the immune system of each patient demands up-to-date therapeutics. Asymptomatic children or those whose illness is moderate without severe hemorrhagic episodes can be monitored and don’t need specific medicinal therapy. In these cases follow-up without medication intervention is the best form of treatment and success of intermittent controls of platelet count until the disease remission. In more severe cases with risks of intracranial, gastro-intestinal and genitourinary hemorrhage, there is the option of platelet transfusion. In cases of chronic IPT, in which no success was achieved with all the attempted therapeutic resources, splenectomy is recommended, despite its performance restricted success.

In the case of acute IPT in children, there are treatment options with corticoids with high doses of intravenous immunoglobulin doses, and more recently with Anti-D immunoglobulin. Intravenous immunoglobulins. The most utilized dose is 50-75µg/kg favors platelet increase within 24 to 48 hours after infusion. Collateral effects verified in the usage of intravenous immunoglobulin are headaches, body tremor, fever, nausea, vomit and aseptic meningitis.

Another treatment option for acute IPT is intravenous immunoglobulins. The most utilized dose is of 0.8 to 1.0/kg/day for 1 or 2 days. In this dosage, it was observed an increase of platelet counts 24 hours after infusion. Collateral effects verified in the usage of intravenous immunoglobulin are headaches, body tremor, fever, nausea, vomit and aseptic meningitis.

Among the alternatives for the treatment of acute IPT, the anti-D immunoglobulin is the most recent option for children who were not submitted to splenectomy and whose erythrocytes are rhesus-positive. The Anti-D mechanism of action, in spite of its incomplete elucidation, acts on the reduction of platelet clearance. The dosage (50-75µg/kg) favors platelet increase within 24 to 48 hours after its administration. The collateral effects observed include headaches, body tremor, fever, nausea, vomiting.
During anamnesis it was verified a history of viral infection before the development of IPT symptoms. In the physical exam, petechiae and hematomas were noticed around the knees (Figure 2) and on the forearm. According to the mother, these purple spots developed spontaneously and/or after minor wounds.

Despite its tendency to be a benign and self-limited condition, acute and chronic IPT are distinct diseases defined by age, platelet counts, bleeding symptoms and previous acute disease before the diagnosis. Regarding buccal health, IPT can frequently bring about buccal mucosa alterations, among which petechiae, ecchymoses and hematomas in areas that are easily traumatized as buccal mucosa, tongue lateral edge and the limit between the hard palate and the soft palate. Other IPT findings include spontaneous gingival bleeding, mucocutaneous bleeding and hemorrhagic vesicles.

This study objective is to provide guidance to the surgeon-dentist with regards to diagnose and attendance of children with Idiopathic Thrombocytopenic Purpura and a clinical case presentation.

CASE REPORT

The patient, 4 years old, was referred to the Centro de Atendimento a Pacientes Especiais (CAPE) at Faculdade de Odontologia da Universidade de São Paulo (FOUSP) with a diagnosis of idiopathic Thrombocytopenic Purpura for odontological follow up. The clinical treatment started after the signature of the Free and Clarified Consent Term for the fulfillment of ethical principles of Helsinki Principle (2000) and Brazilian specific legislation (Figure 1).

Intra-buccal exam revealed no buccal alterations as the result of IPT. Buccal hygiene of the patient was satisfactory and no dental caries and gingival inflammation were observed. An anterior open bite due to pacifier sucking habit, removed 3 months before the first visit, was found. After the clinical and radiographic exam, it was verified the presence of the fusion of elements 72 and 73 (Figure 3).

Full hemogram analysis of the patient showed the presence of thrombocytopenia (78,000/ mm³) and discreet anemia (Figure 4). The patient was not under medication and was attended by a physician to control IPT.
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The treatment consisted of nutrition guidance and buccal hygiene for the patient and her guardians, prophylaxis and topical application of fluoridated varnish (Duraphat®, Colgate). For the correction of the open anterior bite, a removable apparatus with Hawley arch and lingual brace (Figure 5) were used. The patient has been under preventive and orthodontic follow-up for 24 months. She has also been submitted to phonoaudiology treatment to promote the development of correct lingual position (Figure 6).

DISCUSSION

IPT is a common childhood disease with low morbidity and mortality, with or without hemorrhage. Therefore, the surgeon-dentist knowledge with regards to potential alteration caused by this hemopathy is of great importance for dental treatment.

IPT typical signs observed in the patient, as petechiae and hematomas, coincide with the findings in the literature. However, the patient didn't present the usual symptoms of the disease, as the presence of petechiae, ecchymoses and hematomas on the tongue lateral border and on the limit between the hard palate and the soft palate and spontaneous gingival bleeding. Such clinical evidences show IPT attenuation, proven by laboratorial tests.

Medical treatment of patients with IPT assumes different intervention forms in accordance with the clinical conditions observed. Patients affected by IPT in the absence of severe hemorrhage are monitored and don't need specific medicinal therapy, a fact which was verified with this patient.

Regarding dental treatment of children affected by ITP, the clinical performance of the odontopediatrician must be based on previous knowledge of the evolution of the patient illness. Appropriate and comprehensive anamnesis is essential, given that sometimes the patients affected by this pathology don't show buccal clinical signs, as the patient herein described. Hence, the illness might not be noticed by the dentist, if the patient's history is not appropriately reported by their parents.

Dental procedures can be performed in patients with platelet counts above 50,000/mm³. In the need of surgical interventions on a patient whose platelet count is too low, blood transfusion or previous corticosteroid treatment is deemed necessary according to medical prescription. The surgeon-dentist must avoid trauma to the buccal mucosa by carefully puncturing the needles, adapting orthodontic braces and avoiding super-activation. Prescription of antiplatelet drugs, such as acetylsalicylic acid and ibuprofen, must be avoided.

Appropriate control of dental biofilm is of the utmost importance, in order to prevent gingival inflammation and infections. Individual protocols for prevention of caries must be established according to caries risk and age of each patient. If there is the need for restoration treatments, endodontic and surgical treatment, antibiotics prophylaxis can be performed to reduce the risk of postoperative infections.
The treatment of children with IPT must involve an interdisciplinary approach in which the odontopediatrician works together with the hematologist physician.

**CONCLUSION**

Hematological diseases present physical and buccal alterations that are considered important clinical signs for their diagnosis, as buccal mucosa alterations among which petechiae, ecchymoses and hematomas. Notwithstanding, these clinical signs are rare and the professional must be able to recognize Idiopathic Thrombocytopenic Purpura after a critical anamnesis aimed at providing guidance and effective treatment of the patients, as well as refer them to the physician.

**Collaborators**

VF ROSSIER provided clinical attendance to the patient and participated in the article writing. SMCPAC VIEIRA was responsible for the orthodontic assessment and orthodontic attendance and participated in the article writing. AL CIAMPONI was responsible for the conception and drawings of the clinical case and participated of the article writing. RO GUARÊ was in charge of the analysis of clinical data and participated of the article writing.

**REFERENCES**