Acute anemia after dental extraction: A case report

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ABSTRACT

A patient treated with a vitamin K antagonist (VKA) since the implantation of two mechanical heart valves developed acute anemia after the extraction of a tooth. This case report and data in the literature indicate a need for specific measures before, during, and after oral surgery in patients taking anticoagulant therapy: 1) the bleeding risk should be evaluated before the procedure. The INR should be measured routinely, 2) the procedure should be scheduled early in the week to allow an evaluation at the fibrinolysis peak, i.e., 48 to 72 hours after surgery, which is the time of greatest risk of delayed bleeding, 3) the surgical procedure should be appropriate for the elevated bleeding risk, 4) postoperative monitoring is of the most importance, as bleeding is usually delayed in patients on VKA therapy, 5) when poor treatment adherence is expected, day-hospital admission is useful to ensure that the postoperative protocol is implemented properly and to detect early bleeding. The treatment of post-extraction acute anemia includes local hemostasis protocol with the revision of the socket followed by red-blood-cell pack transfusion. A daily fluindione dosage control, and a normal hemoglobin level will allow the patient to leave the hospital.

KEYWORDS

Case Study; Oral Hemorrhage; Acute Anemia; Anti-Coagulant Treatment; Extraction; Hemostasis

1. INTRODUCTION

Few data are available on acute anemia following dental extraction. Most of the reported cases occurred in patients with congenital coagulation disorders [1,2] or in patients receiving thrombosis prophylaxis, usually with vitamin K antagonists (VKAs). We suggest a therapeutic and preventive strategy for VKA-treated patients requiring oral surgery, based on our case of post-extraction acute anemia and on a literature review.

2. CASE REPORT

A 50-year-old woman from North Africa was referred on an emergency basis to the Oral Surgery Unit of the Max Fourestier hospital, Nanterre, France, for extraction of the left maxillary first molar, which was necrotic and exhibited extensive decay of the crown and roots. She had a history of biological valve implantation at 26 years of age for aortic and mitral disease. Degeneration of the biological valves required replacement by mechanical valves when she was 42 years of age. Bleeding from an ulcer in the duodenal bulb during the postoperative period led to partial gastrectomy. Thrombosis prophylaxis treatment with fluindione (Previscan® 20-mg tablets) was initiated, with a target INR of 3 to 4.5.

A few days before her arrival at our department, she had seen a dental surgeon in private practice for pain in tooth #26. The oral surgeon diagnosed pulp necrosis due to mesial decay (Figure 1), opened the pulp cavity, irrigated it with sodium hypochlorite, prescribed amoxicillin (2 g/day for 6 days) given the high risk of endocarditis, and referred the patient to the hospital for extraction of the tooth.

At arrival in our Oral Surgery Unit, the last INR was 3.95, within the therapeutic target range. After discussion with the cardiologist, the surgeon decreased the fluindione dose by 25% to limit the hemorrhagic risk; sche-
dined the procedure for 48 hours later, at the end of the week, with a local hemostasis protocol; and prescribed a single oral dose of 3 g of amoxicillin/clavulanic acid to be taken 1 hour before the procedure [3]. On the day of the procedure, the INR was 3.16, which was within the desired range.

The surgical protocol involved local anesthesia, separation of the roots, removal of the tooth, curettage of the socket, implantation in the socket of absorbable oxidized cellulose (Surgicel®, Johnson & Johnson, Neuchatel, Switzerland) impregnated with 10% tranexamic acid (Exacyl®, sanofi-aventis, France), suturing with absorbable polyglactin (Vicryl rapide® 3.0, Johnson & Johnson), and implantation of Surgicel® mesh secured to the wound with tissue adhesive (Indermil® xfine, Henkel, Ireland).

Local hemostasis was achieved and checked 20 minutes after the end of the procedure. The discharge prescription was for therapeutic amoxicillin/clavulanic acid (2 g/d for 8 days), acetaminophen (40 mg/Kg/d), and gentle mouthwashes with 10% tranexamic acid (Exacyl®, oral solution in vials, 1 g/10 mL) every 6 hours for 5 days. Fluindione (Previscan®) therapy was returned to the usual daily dosage of 3/4 of a tablet (15 mg) on the evening of the procedure, with the agreement of the cardiologist. Written instructions and oral explanations for postoperative care were given to the patient.

At the follow-up visit 4 days later, the patient complained of severe asthenia and reported steady bleeding from the wound, for which she had not seen fit to seek medical advice. Extreme pallor of the skin and mucous membranes was noted. Healing was delayed because of persistent bleeding (Figure 2).

An emergency blood test showed severe anemia with a hemoglobin level of 6.8 g/dL. A mean corpuscular volume of 76 fl (reference range: 80 - 100) and decreased ferritin level (13 µg/L, reference range: 15 - 150) was in accordance with an iron deficiency, probably related to previous bleeding episodes. The INR was abnormally high, at 5.5.

Revision surgery was performed. The blood clot was removed, the socket was revised and filled with Surgicel®, new sutures were performed, and Surgicel® mesh was secured to the wound with tissue adhesive. A resin compression splint was fashioned on an emergency basis and placed in the mouth (Figure 3).

The patient was admitted to the cardiology unit, where she immediately received a red-blood-cell pack. The daily fluindione dosage was decreased by 25%. The blood transfusion was effective, with a hemoglobin level of 8 g/dL.

On the next day, despite the aggressive local hemostasis protocol initiated after the revision procedure, the bleeding recurred. The socket was revised again and filled with Surgicel®, sutures were performed, and a new Surgicel® mesh was secured to the wound with tissue adhesive. The resin compression splint was secured with Coe-Pack® to improve stability (Figure 4).

Two additional red-blood-cell packs were given, which increased the hemoglobin level to 11.8 g/dL. Parenteral...

Figure 1. Initial retroalveolar radiograph.

Figure 2. Bleeding after extraction in a patient taking vitamin K antagonist therapy.
ferric hydroxide-saccharose (Venofer®, Vifor France) was given, followed by oral ferrous sulfate-folic acid (Tardyferon B9®, Pierre Fabre, France).

On the next day, the patient reported feeling better and had no bleeding. Her hemoglobin level was stable at 11.8 g/dL. She was discharged after a total hospital stay of 10 days. Her anticoagulation was stable with a target INR of 3.0 to 4.5. The resin compression splint was removed 10 days after discharge. A clinical and radiological reevaluation 6 months later showed good healing of the bone and mucosa (Figures 5 and 6).

3. DISCUSSIONS

Few data are available on acute anemia following dental extraction. Most of the reported cases occurred in patients with congenital coagulation disorders [1,2] or in patients receiving thrombosis prophylaxis, usually with vitamin K antagonists (VKAs) [4]. Studies of the bleed-
ing risk associated with oral surgery showed that a properly implemented local protocol was effective in ensuring hemostasis during surgery [5,6]. In contrast, an INR value greater than 2.5 was associated with an increased risk of delayed bleeding [7,8], whose prevalence was estimated at about 10% [9]. Available VKAs include the coumarin family (warfarin, acenocoumarol and phenprocoumon), and indandiones (fluindione). Fluindione is frequently used in middle-aged patients in France because of its shorter half-life compared to warfarin (31 h vs 36 - 42 h). Fluindione also differs from warfarin as it is not metabolized by CYP2C9 enzyme of cytochrome P450 system.

In our patient, surgical trauma was minimized, granulation tissue was removed by painstaking curettage, and noncompressive oxidized cellulose (Surgicel®) impregnated with tranexamic acid (Exacyl®) was implanted into the socket. The wound was sutured and covered with Surgicel® mesh secured using tissue adhesive. Examination 20 minutes later showed no bleeding. Thus, the acute anemia was related to delayed bleeding, in keeping with previously published data [6].

Our patient had six risk factors for bleeding. 1) She had a history of major bleeding, consisting in duodenal bleeding after heart surgery. 2) The procedure had to be scheduled on an emergency basis, to alleviate the symptoms, and was performed at the end of the week. She was given a follow-up appointment for the next Tuesday, but the bleeding started over the weekend, when our odontology department was closed. In addition, the patient underestimated the significance of the bleeding and did not comply with the postoperative care instructions, as she did not seek help before her scheduled appointment. 3) Tooth #26 is an antral tooth. The retroalveolar radiograph in our patient clearly showed the proximity of the tooth apex with the floor of the sinus (Figure 1). Disruption of the sinus mucosa can increase the risk of bleeding. 4) VKA therapy is associated with a high risk of postoperative bleeding and, more specifically, with the occurrence of delayed bleeding carrying a risk of potentially severe anemia [6]. In addition, our patient had a very high risk of thrombosis related to the presence of two mechanical heart valves and atrial fibrillation, requiring a high level of anticoagulation with a target INR of 3 to 4.5, compared to 2 - 3 in most other indications for VKA therapy. 5) Antibiotic therapy with amoxicillin-clavulanic acid was started at the end of the surgical procedure. An interaction between this antibiotic and fluindione explains the INR elevation to 5.5, which increased the risk of bleeding [10]. The interaction between VKA and antibiotics mostly results from the destruction of intestinal bacteria producing vitamin K. It is not specific of fluindione but concerns all VKA molecules. Each INR point above 4 doubles the bleeding risk, which was therefore increased 3-fold in our patient [11]. 6) Finally, our patient failed to seek advice before her scheduled appointment, despite persistent bleeding for several days and, consequently, was not able to receive early treatment.

Our case report and data in the literature indicate a need for specific measures before, during, and after oral surgery in patients taking anticoagulant therapy.

First, the bleeding risk should be evaluated before the procedure. An important step is a review of the medical and surgical history including any past bleeding episodes and procedures. The postoperative course after previous surgical procedures provides guidance about the required aggressiveness of the hemostasis protocol. The INR should be measured routinely, and the appropriateness of diminishing the VKA dosage should be discussed with the cardiologist. Finally, the anatomic location of the surgical site is crucial to consider. Computed tomography should be performed if the tooth is near a structure likely to cause bleeding, such as the maxillary sinus, mandibular canal, or tuberosity. If this preoperative evaluation indicates an elevated bleeding risk, an aggressive hemostasis protocol should be used including a hemostatic splint, which can be stabilized during the procedure. In addition, day-hospital admission is indicated when poor treatment adherence is expected and/or the patient lives far from the hospital.

Second, the procedure should be scheduled early in the week to allow an evaluation at the fibrinolysis peak, i.e., 48 to 72 hours after surgery, which is the time of greatest risk of delayed bleeding [12].

Third, the surgical procedure should be appropriate for the elevated bleeding risk. Invasiveness should be minimized to spare the alveolar processes, which ensure stability of the clot. For teeth with multiple roots, root separation is preferable. Sparing the mucogingival junction during full-thickness detachment limits the size of the hematoma. Granulation tissue increases the risk of surgical site infection and bleeding and should be completely removed by careful curettage of the socket. The full hemostatic protocol includes noncompressive implantation of hemostatic material into the socket, effective suturing, oxidized cellulose secured over the wound, and use of a hemostatic splint.

Fourth, postoperative monitoring is of the utmost importance, as bleeding is usually delayed in patients on VKA therapy. The postoperative treatments must be described in detail and explained to the patient, family, and healthcare professionals (e.g., nurse or nursing assistant). An antifibrinolytic agent (tranexamic acid, Exacyl®) is used as gentle mouthwashes three times daily for 5 days [5] and as compression packs in the event of bleeding [13-15]. Selection of the antibiotics and analgesics should take into account the many interactions between
these two families of drugs and VKAs. In patients on VKA therapy, nonsteroidal antiinflammatory agents should be avoided and administration of amoxicillin/clavulanic acid, similar to all other antibiotics, requires close monitoring of INR values [10,16-18]. Alternatively, amoxicillin alone or pristinamycin can be used. Detailed instructions about postoperative care should be given to the patient orally and in writing. When there is a language barrier, help should be sought from an interpreter, for instance a family member. Optimal treatment adherence is crucial to successful treatment and can be obtained only if the patient understands the postoperative regimen and its importance. An emergency telephone number should be communicated to the patient.

Fifth, when poor treatment adherence is expected, day-hospital admission is useful to ensure that the postoperative protocol is implemented properly and to detect early bleeding. Strict compliance with these rules should increase the success rate of oral surgery procedures by avoiding complications that may be threatening and/or induce severe quality-of-life impairments.

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REFERENCES


