Early complication after total knee arthroplasty in a haemophilia A patient

Wczesne powikłanie po endoprotezoplastyce stawu kolanowego u chorego na ciężką hemofilię A

Pawel Ambroziak¹, Piotr Żbikowski¹, Andrzej Kotela¹, Jerzy Windyga², Ireneusz Kotela¹,³

¹Orthopedic and Trauma Clinic, Central Clinical Hospital of the Ministry of Interior, Warsaw, Poland
Head of Department: Prof. Ireneusz Kotela MD, PhD
²Department of Disorders of Hemostasis and Internal Medicine, Institute of Hematology and Transfusion Medicine, Warsaw, Poland
Head of Department: Prof. Jerzy Indyga MD, PhD
³Faculty of Sciences, Jan Kochanowski University, Kielce, Poland
Head of Department: Prof. Ireneusz Kotela MD, PhD

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Abstract

Total knee replacement in patients with haemophilia A is a challenging procedure with high risk of complications. Due to the massive destruction of the joint and significantly reduced range of motion, total knee replacement should be performed only by a surgeon with high degree of expertise and experience. During the perioperative period, patients require factor VIII (FVIII) replacement therapy supervised by a haematologist, under control of plasma activity levels. Possible early complications include delayed wound healing, soft tissue and joint bleeding, development of pseudoaneurysm and early infection. Once complications occur, prompt detection and introduction of proper treatment is fundamental.

Streszczenie

Endoprotezoplastyka całkowita stawów u chorych na hemofilię A jest operacją trudną i obarczoną dużym ryzykiem wystąpienia powikłań. Z powodu masowych zmian zwyrodnieniowych i często znacznego ograniczenia zakresu ruchu przed operacją implantacja endoprotezy wymaga dużego doświadczenia operacyjnego. W okresie okołooperacyjnym należy pacjentowi podać koncentrat brakującego czynnika krzepnięcia VIII (factor VIII – FVIII) pod kontrolą jego aktywności w osoczu. Leczenie substytucyjne powinno przebiegać pod ścisłym nadzorem hematologa. Do możliwych wczesnych powikłań operacji należą opóźnione lub zaburzone gojenie rany, krwawienie do tkanek miękkich lub stawu, rozwój tętniaka rzekomoego, wcześniak infekcja. Wystąpienie jednego z wyżej wymienionych powikłań wymaga wczesnego rozpoznania i odpowiedniego leczenia.

Introduction

Frequent, recurrent joint bleeds in patients with severe haemophilia A (factor VIII plasma activity levels < 1 IU/dl) lead to destruction of the joint and rapid development of massive degenerative changes: haemophilic arthropathy. These changes affect not only the articular surface and bone but also the soft tissues of the joint. Synovial fibrosis due to chronic inflammatory changes and a tendency towards antalgic limb positioning cause gradual limitation of range of motion (ROM) – flexion contracture. Reduction of weight bearing of the affected limb leads to muscle atrophy. These changes occur commonly in knee joints, ankles and elbows. Degenerative changes as a result of joint bleeds lead to significant disability [1]. In cases of insufficient replacement of deficient clotting factor, advanced arthropathy may be present in the age range 15–25 years. Patients may often need to use crutches or a wheelchair for locomotion. When arthropathy progresses despite treatment and physiotherapy, the only effective way to treat it is total joint arthroplasty [1–3].

Despite specific, precise guidelines for the treatment of patients with haemophilic arthropathy, total knee arthroplasty is performed rarely and only at specialised centres. The problems usually encountered include the lack of clear orthopaedic recommendations, significantly greater risk of complications and difficulties in creating an experienced team (including haematologist, orthopaedic surgeon, anaesthetist and
physiotherapist). As well as typical risks and potential complications of joint arthroplasty, complications associated with blood clotting can occur. Surgical difficulties are associated with limitations of ROM and abnormal joint and limb axis. Arthropathy often involves more than one joint; hence, proper planning of the sequence of surgical procedures is the key to success. The recommended sequence is to perform hip replacement first, then knee arthroplasty. Choice of the proper implant is also of paramount importance. Stable implant placement often requires additional bone grafting. Implant survival time may be shorter because of weaker bonding to osteoporotic bone. A higher risk of aseptic or septic implant loosening is noticed. Early wound healing problems are another possible complication. To perform this kind of surgical procedure safely it is necessary to create an experienced team to supervise the patient during treatment. Access to a suitably equipped, specialised laboratory is mandatory.

**Case report**

A 25-year-old patient suffering from knee arthropathy caused by repetitive joint bleeds in the course of severe haemophilia A was admitted to the Department of Haemostatic Disorders and Internal Medicine of the Institute of Haematology and Transfusion Medicine in Warsaw for preparation for elective total knee replacement. Comorbidities were epilepsy and chronic hepatitis C virus infection (HCV). Patient height was 178 cm, weight 80 kg and body mass index (BMI) 25 kg/m². Advanced knee arthropathy was associated with severe ROM restriction and pain. Range of motion before surgery: extension deficit 10°, flexion 90°. The joint was stable in the frontal and sagittal planes with correct axis of the limb (Figure 1).

Basic preoperative laboratory tests were correct except for a highly elongated activated partial thromboplastin time (APTT) due to the absence of factor VIII (< 1 IU/dl). The most important laboratory test allowing the patient to elective surgical procedure is the test for the presence of factor VIII inhibitor. In the discussed patient, inhibitor of factor VIII was excluded, so there were no haematological contraindications to elective surgery. Two hours before surgery, factor VIII concentrate was administered at a dose of 50 IU/kg [1]. Factor VIII plasma activity 30 min after injection of the concentrate was 104 IU/dl (normal range 50–150 IU/dl); therefore, inherited deficiency of factor VIII was corrected.

The patient was transferred to the Orthopaedic and Trauma Clinic of the Central Hospital of the Ministry of the Interior to perform total knee arthroplasty. The surgery was performed under general anaesthesia in a typical manner (Figure 2). During the operation patella maltracking was noticed and lateral retinaculum was released. Before closing the wound the pneumatic tourniquet was released and meticulous haemostasis was made. Two drains were left in the wound. In the perioperative period (I–II week) factor VIII plasma activity was maintained at a level above 80 U/dl, and was checked every 24 h [4]. Thromboprophylaxis was conducted using nadroparin 0.6 ml subcutaneously every 24 h. For the first 4 days, wound bleeding was noticed with no bleeding into the drains. The drains were removed on the second postoperative day.

Physiotherapy was postponed until day 4, when wound bleeding had stopped. In the following days the correct course of treatment was noticed. An considerable swelling of the knee was present. Narcotic drugs (morphine) were used for analgesia. Non-steroidal, anti-inflammatory drugs were not used due to the potential impact on blood coagulation parameters.

On the 15th day after the surgery slight wound bleeding occurred again, despite maintaining proper FVIII plasma activity (80 U/dl); therefore, there was no indication for modification of substitution of FVIII [4]. Bleeding was accompanied by increasing...
Continuation of physiotherapy was recommended. Wound revision in good condition, with 0–70° ROM of the operated joint and without swelling or pain. Continuation of the physiotherapy was recommended. Since the wound was healed, physiotherapy was carried out with FVIII administration 3 times a week. The patient remains under outpatient observation.

Discussion

Replacement of deficient clotting factor is necessary when carrying out orthopaedic operations in haemophilia. During the perioperative period, patients receive FVIII concentrate intravenously, maintaining its plasma activity at the level of 100 U/dl. Treatment progress after wound revision was typical. The drain was removed on the 3rd day with a total blood loss of 50 ml. Intensive physiotherapy was reintroduced, including walking on two crutches with partial weight bearing and ROM exercises. Sutures were removed on the 12th day. The patient was discharged on the 18th day after surgical wound revision in good condition, with 0–70° ROM of the operated joint and without swelling or pain. Continuation of the physiotherapy was recommended. Since the wound was healed, physiotherapy was carried out with FVIII administration 3 times a week. The patient remains under outpatient observation.

Arterial bleeding requires inspection of the wound and its careful closure. An additional, and very useful, method can be angiography with embolisation of the injured vessel [7–9]. In the case described above, active arterial bleeding was revealed during haematoma removal. After exclusion of possible haematological causes of bleeding (proper factor VIII plasma activity, excluding the presence of inhibitor), surgical wound inspection was performed without additional imaging (ultrasound, angiography). It can be stated that the bleeding was the result of an intraoperative vascular injury.

Arterial bleeding in a patient with haemophilia A can occur even with normal plasma activity of clotting factor. Further increases in factor concentrate doses are usually useless and may lead to thrombotic complications. In such cases, surgical intervention and wound inspection is required. There are few reports on complications in the form of a postoperative pseudoaneurysm or persistent joint or soft tissue haematoma [7, 10–15]. The recommended procedure is arteriography with embolisation of the injured vessel or surgical wound inspection and closure of the injured vessel under direct vision.

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Address for correspondence:

Paweł Ambroziak
Orthopedic and Trauma Clinic
Central Clinical Hospital
of the Ministry of Interior
ul. Wołoska 137, 01-912 Warsaw, Poland
Phone: +48 22 508 13 70
E-mail: pawel.ambroziak@wp.pl