Case Report

Compartment Syndrome, Femoral Nerve Lesion and Recurrent Haemarthrosis in one and the same Patient after Total Knee Arthroplasty

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Abstract

We present a case of a 53 year old patient that underwent total knee arthroplasty in our hospital for degenerative arthritis and developed a series of very uncommon complications. Compartment syndrome of the calf occurred postoperatively as a result of bleeding from a large, ruptured knee joint cyst. Femoral nerve lesion was diagnosed and was most likely the result of traumatic migration of the femoral nerve catheter in the nerve sheet. Finally, our patient suffered from recurrent haemarthrosis in which anticoagulant therapy is thought to have played a key role.

This series of very uncommon complications occurring in one and the same patient stresses the importance for orthopaedic
surgeons to continuously and carefully monitor their patients postoperatively in order to adequately diagnose complications, also the rare ones, and to take appropriate measurements to minimize dramatic outcomes of these complications.

**Keywords:** Compartment syndrome, femoral nerve lesion, recurrent haemarthrosis, total knee arthroplasty.
Introduction

Total knee arthroplasty (TKA) is one of the most successful operative procedures and survivorship figures of 98% at 15 years have been published (Ritter et al., 2001). Serious complications after joint arthroplasty are rare but do occur at a combined rate of 3% (Pulido et al., 2008). Deep venous thrombosis (DVT) with the potential to propagate a potentially lethal pulmonary embolus (PE) is the most feared early complication of TKA and the prevalence of asymptomatic DVT without thromboprophylaxis ranges from 41% to 85% when examined by venography at 7 to 14 days (Colwell, 2009). Use of effective prophylactic agents can reduce the rate of postoperative DVT by 60% to 70% (Geerts et al., 2008). Prevalence of PE ranges from 0.9% to 28% for all PE and 0.1% to 2% for fatal PE (Colwell,
Furthermore, infection is a feared and serious complication. The rate of prosthetic joint infection is currently estimated to be between 1.0 and 3.0% (Peel et al.). Peroneal nerve palsy and recurrent haemarthrosis are uncommon complications. The rate of peroneal nerve palsy has been reported to be 0.6% (Idusuyi and Morrey, 1996) and the rate of recurrent haemarthrosis has been reported in up to 1.6% of patients following TKA (Saksena et al., 2010). We report on a case of a 53 year old woman who developed a cascade of uncommon but major complications following total knee arthroplasty that were related to both surgery and anaesthesia.
Case

Late August 2009, a 53 year old patient came to our hospital with complaints related to severe traumatic arthritis of the left knee. She suffered a ski trauma in 1987 and had undergone several arthroscopic procedures to improve knee function. Her primary complaints were of a painful, frequently swollen knee. Physical examination revealed a varus axis of 6 degrees, a flexion of 90 degrees and an extension deficit of 5 degrees. Standard radiographs showed marked arthritis of all 3 compartments of the knee. Patient was otherwise healthy and did not use any medication.

Preoperative standard workup with laboratory testing’s and urine analysis was normal. Patient was scheduled to undergo
total knee replacement in April 2010. Operative procedure was performed under spinal anaesthesia. A tourniquet was used only during cementation of the TKA components. It was inflated to 250 mmHg for 20 minutes. A standard medial parapatellar approach was used and a standard, cemented, cruciate retaining prosthesis (Vanguard™ Complete Knee System, Biomet, Inc., Warsaw, IN) was inserted. Patella resurfacing was considered unnecessary. Ligament release was not necessary. A retransfusion drain (Bellovac ABT, Astratech, Mölndal, Sweden) was used. Total procedure time was 60 minutes and no complications occurred intraoperatively with either spinal anaesthesia or surgery itself. A femoral nerve catheter (Contiplex, B. Braun Medical) was inserted uncomplicated and painless after ultrasound localisation and using nerve stimulation technique. Patient started with Fondaparinux 5mg/ml, 0.5 ml
(GlaxoSmithKline) 6 hours post-surgery to prevent tromboembolic complications.

On the first day postoperative, a pulling force was exerted on the femoral catheter by accident and from that moment on the patient reported a sharp pain and a cold sensation every time fluid was injected into the catheter. On the second postoperative day, after bandage removal and after removal of the femoral catheter, patient started to complain of a painful and seriously swollen calf. Ultrasound was performed to rule out deep vein thrombosis and showed an open venous system up from the popliteal vein. However, spontaneous compression of the popliteal vein itself was seen at the level of the calf due to diffuse swelling. A Bakers cyst was noted measuring 4 centimetres in diameter. At this time, there were no signs of a compartment
syndrome. A day later, however, physical examination revealed a 50% loss of strength of the anterior tibial muscle and the extensor hallucis longus muscle and loss of sensation in the first web space, in combination with extremely painful passive mobilisation of the forefoot. A few hours later, spontaneous bleeding in the upper leg and the knee joint occurred. Angiography was performed but no damage of the arterial structures could be diagnosed (figure 1). Emergency fasciotomy of the anterior, lateral and superficial and deep compartments of the lower leg and arthrotomy of the knee were performed for decompression under general anaesthesia. A tourniquet was not used. 200 cc of blood was drained from the knee joint. In the superficial flexor loge a large haematoma between gastrocnemius and soleus muscle was found, resulting from bleeding from a
large, ruptured knee joint cyst. This extensive haematoma caused compression of all compartments in the lower leg.

Figure 1: Arteriography Showing no Evidence of Arterial Injury or a Pseudoaneurysm
Signs of the compartment syndrome disappeared completely during the following two days after fasciotomy, but at this time weakness of the quadriceps muscle was noted. Analysis by the neurologist revealed a paralysis of the quadriceps muscle. There were no signs of altered skin sensibility. Ultrasonography of the groin excluded the presence of a local hematoma and no signs of compression of the neurovascular structures in the groin were seen. Electromyography performed 2 weeks later revealed a lesion of the femoral nerve with only minimal spontaneous recovery in the vastus lateralis muscle. The patient stayed admitted. Six weeks after initial surgery, patient complained of a painful, swollen knee. Punction of the knee revealed a new haemarthrosis and three days later it was decided to debride the knee arthroscopically because of persistent pain. This procedure was performed under general anaesthesia and again no
tourniquet was used. Blood thrombi were evacuated from the knee joint. There were no signs of impingement or proliferative synovitis. Anticoagulant therapy was stopped at this time. No new episodes of bleeding occurred after this intervention. Analysis for coagulation disorders was performed but no abnormalities could be detected. Two weeks later, a new elektromyography was performed to re-evaluate the femoral nerve paralysis. No signs of recovery could be observed at this stage and prognosis was uncertain.

The patient was eventually discharged two days later, nearly 2 months after initial surgery and she was registered for intensive rehabilitation. Seven months after the initial surgery, weakness of the quadriceps muscle was still remarkable. A new electromyography showed a normal conduction in the femoral
nerve and signs of reinnervation of the quadriceps muscle. No haemarthros developed since the day of discharge. Flexion of the knee was only 45 degrees and a 5 degree extension deficit remained. At this moment patient was bound to a wheelchair for most of the time. At 10 months post-surgery, residual weakness of the quadriceps muscle was limited and knee function had improved to 70 degrees of flexion and a 5 degree extension deficit.

Discussion

Signs and symptoms of compartment syndrome include: pain out of proportion to the injury and worsening of the pain with passive stretching of the involved muscles, swelling and coldness. Late signs are paraesthesia, pulselessness and paralysis, although
the presence of a pulse cannot rule out the presence of compartment syndrome (Sloper et al., 2003). Compartment syndrome complicating knee arthroscopy is documented (Kim et al., 2002, Mont et al., 1996), but it is only rare after total joint replacement and a specific cause in these cases cannot always be identified (Hailer et al., 2007). Some reports are published on compartment syndrome after TKA. Tang and Chiu were the first to describe occurrence of compartment syndrome of the calf after early postoperative dislocation of a TKA (Tang and Chiu, 2000). Haggis et al. reported on 7 cases of compartment syndrome following TKA and identified risk factors for the development of a compartment syndrome (Haggis et al., 2006). These factors include: soft-tissue compromise, often related to previous surgery, injury or complex surgery and vascular compromise or intra-operative vascular injury. In our case, no such risk factors
were present. The compartment syndrome was the result of bleeding from a knee joint cyst. This has been reported previously to be one of the causes of compartment syndrome (Haggis et al., 2006). Bleeding did not occur directly in the anterior compartment, but compression of this compartment was probably indirect due to a haematoma in the flexor compartment with subsequent spontaneous compression of the popliteal vein. We did not perform pressure measurements but other causes for peroneal nerve involvement were unlikely because of normal peroneal nerve function directly postoperative and the absence of sonographic abnormalities.

An overall incidence of temporary femoral nerve injury of 0.6% has been reported after femoral catheter placement with the use of electrical stimulation guidance (Feibel et al., 2009). Incidence
of permanent neurologic injury, as manifested by weak quadriceps power, diminished sensation, and dysesthesia, is estimated to be 0.2% (Feibel et al., 2009). Femoral neuropathy can be caused by iliacus hematoma and is a well-known complication in hemophiliacs or in patients undergoing anticoagulant therapy (Puechal et al., 1992, Rodriguez-Merchan, 2012). In patients without coagulation disturbances, however, femoral neuropathy caused by iliacus hematoma rarely occurs, the cause is usually traumatism instead (Puechal et al., 1992). Concerning these traumatisms, it is hypothesised that the potential for coiling of the perineural catheter and pressure injury to the nerve, rather than intraneural puncture, pertains a risk of neurologic injury (Feibel et al., 2009). A similar trauma mechanism to the femoral nerve has probably taken place in our case, where a pulling force had been exerted onto the catheter
with secondary movement of the catheter in the neural sheet. From this moment on, patient reported a sharp pain, every time fluid was injected into the catheter. Pain or paresthesias on injection of local anesthetics in patients who had nerve damage have been observed by others (Auroy et al., 1997, Auroy et al., 2002). It is possible that these injections have further aggravated the damage to the femoral nerve by increasing intraneural pressure.

Nerve palsy can be a complication of tourniquet use and a total tourniquet time of more than 120 minutes is a significant risk factor for the development of peroneal nerve palsies. There are, as far as we know, no femoral nerve laesions reported in current literature that were the result of long tourniquet time. Given the absence of these reports, the obvious trauma exerted onto the
catheter in our case and a tourniquet time of only 20 minutes, it is unlikely that the use of a tourniquet has played a role in the pathogenesis of the femoral nerve injury in our case. Time to neurologic recovery of a femoral nerve lesion ranges from 36 hours to 6 months (Feibel et al., 2009, Capdevila et al., 2005). In our case, at 7-month follow-up, clinical weakness of the quadriceps muscle persisted, but EMG showed evidence of reinnervation (fasciculation potentials). The long-term prognosis therefore seems favourable.

Several causes for recurrent haemarthrosis have been pointed out. Early bleeding is mostly due to direct vascular injury, arteriovenous fistula, arterial aneurysm and pseudoaneurysm (Saksena et al.). Furthermore, early bleeding may be the result of a loose tag of synovial tissue remaining from surgery that
becomes entrapped between the moving components of the prosthesis early in the postoperative course (Kindsfater and Scott, 1995). It also can be the result of a platelet factor 3 availability defect (Malhotra et al., 2005). In our case, no such cause for recurrent bleeding could be identified. It has been stated previously that causes of haemarthrosis remain unclear in 35,8% of cases (Suzuki et al., 2006). Unexplained haemarthrosis after TKA, as was the case in our patient, should ideally be treated in stages (Rukavina et al.). In every case needle aspiration should be performed first for relief of pain and stiffness, followed by cooling, elevation, rest and discontinuing anticoagulants when possible (Ohdera et al., 2004). If conservative treatment is not successful a femoral arteriography should be considered with the possibility for immediate embolization of the source of bleeding. If embolization is unsuccessful, arthroscopy is the next step.
Entrapped tissue and any source of bleeding can then be localized and treated. If this approach is unsuccessful an open synovectomy is indicated (Rukavina et al.). In our case, bleeding stopped after discontinuing anticoagulant therapy. However, this was only done at a later stage, after invasive therapies had already been performed. Theoretically, these invasive therapies could have been unnecessary in our case if anticoagulants were discontinued in an early phase. Moreover, the use of fondaparinux 5mg/ml on itself could have been an initiating factor for the bleeding episodes as it is stated that the risk of major bleeding (defined as fatal bleeding, bleeding involving a critical organ, bleeding leading to reoperation and overt bleeding with a bleeding index of ≥ 2) is significantly higher than with enoxaparin 30mg twice daily (Bauer et al., 2001, Turpie et al., 2002). Furthermore, a statistically significant relationship was
reported between the incidence of major bleeding and the timing of the first fondaparinux injection between 3 and 9 hours post-surgery (Turpie et al., 2002). In our case, injection was administered 6 hours after surgery. Given these observations, the use of fondaparinux as anticoagulant therapy has therefore likely played a role in the pathogenesis of this bleeding complication.

**Conclusion**

Total knee arthroplasty is one of the most successful procedures with long survival and is perceived as a save procedure. In this case three very uncommon complications occurred together in one patient. We believe this case stresses the importance for orthopaedic surgeons to continuously and carefully monitor their patients postoperatively in order to adequately diagnose complications, also the rare ones, and to take appropriate
measurements to minimize dramatic outcomes of these complications. Furthermore, we believe that careful and critical use of anticoagulants is mandatory for lowering the risk of postoperative bleeding complications in patients with TKA.

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