Prophylaxis of venous thromboembolism in knee replacement surgery

Carlo Rostagno¹, Christian Carulli², Alessandro Cartei², Claudia Ranalli¹, Massimo Curcio¹, Alessandra Cammilli¹, Gian Luca Polidori¹, Massimo Innocenti²

ABSTRACT

Venous thromboembolism (VTE) is still a frequent and sometimes severe complication after knee replacement surgery. Both pharmacological and non-pharmacological measures have been widely used to decrease VTE risk. Non pharmacological treatment include measures directed to decrease the effects of blood stasis, intermittent pneumatic compression device (IPCD) and graduated compression stockings, and mechanical devices. Pharmacological prophylaxis of venous thromboembolism (VTE) is associated with a more significant decrease in the incidence of deep venous thrombosis (DVT) and related complications after knee arthroplasty however anticoagulation may increase the risk of postoperative bleeding and related complications, in particular the need for re-intervention. Aim of present review was to suggest practical approach to DVT prophylaxis in patients undergoing knee arthroplasty.

INTRODUCTION

Despite widespread use of antithrombotic prophylaxis the risk of deep venous thrombosis (DVT) after major orthopedic surgery is still not negligible [1,2]. DVT in these patients is frequently distal, asymptomatic and diagnosed only by routine ultrasonography. Usually the thrombus lyses with complete recovery. Late pulmonary embolism however has been reported even 40 days after surgery [3-5]. At present no clear indication about the need for prolonged anticoagulation in small distal DVT has been provided.

In an observational study Wang et al [6] reported that 160 out of 359 patients developed distal DVT after TKR. Eighty three (52%) involved the gastrocnemius and soleus muscular veins; 38 (46%) were isolated muscular DVT and 45 (54%) involved muscular branches and major leg veins. Rates of clinical symptoms, late DVT, thrombus propagation and pulmonary embolism in patients with isolated muscular DVT were comparable between patients with isolated muscular DVT in comparison to patients with DVT in the major leg veins and patients with combined lesions [7].

The incidence of symptomatic DVT is comprised between 1.5-10% of patients undergoing knee arthroplasty, while fatal pulmonary embolism is relatively uncommon (0.5 to 1.5%) [1,8]. These data are confirmed by a systematic review that reported an incidence of VTE of 1.09% for knee surgery (DVT 0.63%, pulmonary embolism –PE- 0.27%) in patients in adequate drug prophylaxis [8]. A recent paper from Waitemata District Health Board – New Zealand – reports in a 4 year period a cumulative incidence of symptomatic VTE within 90 days of surgery of 3.29% [9]. The median time from surgery to diagnosis was 7 days. Deep vein thrombosis comprised 75% of cases, 77.6% distal and 23.2% proximal. Pulmonary embolism comprised 26.5% of VTE; 47.7% had right heart strain on computed tomography/echocardiography. Of patients developing VTE, 85.5% had pharmacological thrombo prophylaxis - aspirin 73%, LMWH 20 mg 16%, LMWH 40 mg 16%, therapeutic LMWH 3%, unfractionated heparin twice daily 1%, and warfarin 4%; 75.6% received mechanical prophylaxis, while 4% of patients received no prophylaxis.

The main limitation of pharmacologic VTE prophylaxis is the risk of bleeding and related complications, in particular the need for re intervention. Major bleeding following TKR has been estimated between 1 and 3 percent. These data however suffer from the limitation of the variability in the definitions of bleeding used in published literature and few data in control patients.

PATHOPHYSIOLOGY

Knee replacement surgery is associated with trauma to the soft tissues and bone with release of cytokines and other pro-coagulant factors [10]. Excessive release of tissue factor from injury to the vascular wall leads to generation of large amounts of thrombin, which favors platelet activation and thrombus formation [11].
Finally blood venous stasis occurs as a result of several different factors: immobilization, edema, poor calf muscle pump and mechanical haemostatic devices used in surgery room. Manipulation of the limb and the use of a tourniquet in patients undergoing TKA also contribute to venous stasis. [12]. VTE risk is highest in the first days after surgery, on average 7 days after TKA in comparison to 17 days after THA [1]. Early discontinuation of pharmacological prophylaxis favors extension of preexistent asymptomatic thrombosis or development of new thrombotic events.

The risk to develop DVT is significantly higher in patients undergoing knee replacement in comparison to both hip replacement and treatment of hip fracture [13-14] with a 4:1 ratio between distal and proximal thrombosis. Age, underlying pre-existing diseases (cancer, obesity, previous DVT, thromboprophylactic conditions including those genetically determined), injury related factors and finally factors related to medical procedures significantly influence DVT risk [15].

Female sex and previous DVT were the more relevant risk factors for DVT independently from the drug used for pharmacological prophylaxis in the ORTHO-TEP study [16] . In a study by Kang et al [17] DVT was detected by B-mode ultrasonography performed 7 days after TKR in 175/1025 patients aged >75 years. Diabetes (P = 0.014), high body mass index (BMI) (P = 0.003) and hyperhomocysteinemia (P < 0.001) were significantly more frequent in the thrombus group. A significantly greater proportion of patients in the non-thrombus group had early postoperative activity (P < 0.001) and used a foot pump (P < 0.001). Operative duration was significantly longer in the thrombus group (P = 0.012). DVT was more frequent in bilateral than unilateral knee arthroplasty (P < 0.01). Multivariate logistic analysis revealed BMI, long operative duration, bilateral knee arthroplasty, and time to the activity after the operation to be predictive factors of DVT. At 6-month follow-up in DVT group, 4.7% of patients had pulmonary embolism and 18.8% had recurrent DVT.

**ProhPylaxis of DVT –PE**

In patients undergoing major orthopaedic surgery pharmacological and non-pharmacological measures are available for VTE prophylaxis. At present parenteral agent, low dose unfractionated heparin (UH), low molecular weight heparins (LMWH) and fondaparinux are mainly used for drug prophylaxis. Direct oral anticoagulants may be indicated in elective surgery in patients with expected poor adherence to parenteral route. Non pharmacological treatment include measures directed to decrease the effects of blood stasis, intermittent pneumatic compression device (IPCD) and graduated compression stockings, and mechanical devices, IVC filters to avoid migration of eventual emboli to pulmonary circulation. IPCD and graduated compression stockings anyway should be considered as adjunctive measures or used when contraindications exist to drug prophylaxis.

**Low dose unfractionated heparin**

Low dose subcutaneous unfractionated heparin (UH) has been widely used for VTE prophylaxis before 1990. At present indication for low dose unfractionated heparin are limited, as clear evidence has been provided of superiority of LMWH [18]. In advanced renal failure (creatinine clearance < 30 ml/min) UH may instead of LMWH in order to decrease haemorrhagic risk.

**Low molecular weight heparin (LMWH)**

LMWH at present are the drugs of choice for pharmacological prophylaxis of DVT in orthopaedic surgery [19-21]. After TKR treatment decreases by more than 50% the risk of post-operative asymptomatic DVT in comparison to no prophylaxis. Several studies have clearly demonstrated the superiority of LMWH in comparison to UH and adjusted dose warfarin with a similar, or superior in the case ofwarfarin, safety profile [22-24]. The risk for heparin induced thrombocytopenia (HIT) is lower with LMWH than with UH [25]. Effects on symptomatic DVT or PE were not significantly different when LMWH was compared to fondaparinux, however the latter showed an higher bleeding rate leading to re-intervention [26-29]. Each LMWH has a specific molecular weight distribution that influence anticoagulant activity, duration of action, and renal clearance, so each agent must be considered a unique drug and one product cannot always be substituted for another (table 1).

**Table 1. LMWHs available for DVT prophylaxis**

<table>
<thead>
<tr>
<th>Active principle</th>
<th>Standard dosage</th>
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<tbody>
<tr>
<td>Dalteparin</td>
<td>5000 IU before surgery, thereafter 5000 IU /day</td>
</tr>
<tr>
<td>Enoxaparin</td>
<td>40 mg (4000 IU) 12 h before surgery, thereafter 40 mg daily (european dose) , 30 mg (3000 IU) before surgery, thereafter 30 mg b.i.d</td>
</tr>
<tr>
<td>Nadroparin</td>
<td>38 IU /kg 12 h before surgery and 12 hours after. 38 IU /kg for the next 3 days, then increase the dose to 57 IU/day</td>
</tr>
<tr>
<td>Reviparin</td>
<td>4200 IU before surgery, thereafter 4200 IU /day</td>
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At present no clear evidence has been provided on the optimal dose to be employed nor regarding efficacy difference among different molecules. For enoxaparin European approved regimen is 40 mg (4000 IU) once –daily, US regimen is 30 mg b.i.d. Dose for patients with decreased renal function must be adjusted in relation to creatinine-clearance. Duration of treatment, initially limited to 10-14 days, after the demonstration of a non negligible risk of DVT after hospital discharge has been extended over 30 days [30].

**Fondaparinux**

Fondaparinux, a synthetic drug, indirectly inhibits activated factor X through an high affinity binding with antithrombin. Only one study examined the effects of fondaparinux in comparison to placebo [31]. Although a
decrease of PE was reported, other end-points were not significantly affected and, due to increased bleeding risk, it was impossible to assess a beneficial effect on overall mortality. Results from 4 randomized studies comparing fondaparinux to enoxaparin were reported by Turpie et al [32]. The original primary efficacy endpoint consisted of a composite of deep-vein thrombosis detected by mandatory bilateral venography, documented symptomatic deep-vein thrombosis, or pulmonary embolism up to day 11. The incidence of VTE was 13.7% (371 of 2,705 patients) in the enoxaparin group compared with 6.8% (182 of 2,682 patients) in the fondaparinux group, with a risk reduction of 55.2% in favour of fondaparinux.

ORTHODO-TEP registry compared the efficacy and safety of fondaparinux and LMWH in 3819 patients undergoing major orthopaedic surgery [16]. The main limitation of the study was its retrospective nature and the comparison of two different periods, the first in which LMWH (3000-6000 aX U) and in the second with fondaparinux 2.5 mg used as thromboprophylaxis. Symptomatic DVT occurred in 4.1% of patients treated with LMWH in comparison to 5.6% in patients treated with fondaparinux. No significant difference was found between the two regimens in the incidence of pulmonary embolism and death. Despite similar overall incidence of bleeding the need for surgical revision was lower with fondaparinux. At multivariate analysis previous VTE and female sex were associated with an increased risk of DTV.

**Vitamin K antagonists**

In United States adjusted dose warfarin is still used for DVT prophylaxis in orthopaedic surgery [33-34]. Extended vitamin K antagonists treatment (>4 weeks) after THA was associated with a 4.6% absolute decrease of VTE with a small increase in the risk of bleeding [35].

**Aspirin**

Studies on the effects of aspirin in DVT prophylaxis failed to demonstrate a favourable effect in patients undergoing THA, TKA or finally treatment for hip fracture [36-37]. It must be emphasized that serious methodological limitations impair a correct interpretation of these studies. Raphael et al [38] analyzed retrospectively a total of 28,923 patients underwent THA or TKA between January 2000 and June 2012. Patients were treated with either aspirin (325 mg twice daily; 2800 patients) or warfarin (26,123 patients). The overall symptomatic PE rate was lower (p < 0.001) in patients receiving aspirin (0.14%) than in the patients receiving warfarin (1.07%). This difference did not change after matching. The aspirin group also had significantly fewer symptomatic DVTs and wound-related problems. Jiang et al [39] randomized 120 patients undergoing TKA to aspirin combined with mechanical measures postoperatively (group A), and respectively low-molecular-weight heparin (LMWH) sodium and rivaroxaban sequentially in combination with mechanical measures (group B). Postoperatively DVT was detected in 10 of 60 patients in group A (16.7%) compared with 11 of 60 in group B (18.3%) (P = 0.500). There were no cases of symptomatic VTE or death during the follow-up period. Patients in group A had the lower blood loss index as compared with patients in group B. The cost of VTE prevention analysis indicated a cost reduction using aspirin in group A compared with using LMWH and rivaroxaban in group B.

**Direct oral anticoagulants**

In the last decade several oral direct anticoagulant have been introduced in clinical practice. These agents have predictable anticoagulant effect, limited drug or food interactions and therefore may be administered at fixed doses without the need for regular laboratory monitoring and dose adjustments. Dabigatran exilate is a selective thrombin inhibitor while apixaban, rivaroxaban and edoxaban are direct Xa factor inhibitors (figure 1). These drugs differ for pharmacological and pharmacodynamics characteristics. Clinical studies, usually, have compared these drugs in TKA against enoxaparin at US or Europe dosages (respectively 3000 IU -50 mg – bi.d. and 4000 IU daily). Most published trials have a similar primary end point e.g. combined symptomatic or asymptomatic (diagnosed by venography) DVT, non fatal pulmonary embolism and all cause mortality. A higher compliance after hospital discharge has been reported in comparison to low molecular weight heparin. Potential safety concerns must to be cleared by time. Dabigatran is a substrate for efflux transporter P-gp: administration in association with a P-gp inhibitor (amiodarone, verapamil, quinidine, ketoconazole, dronedaron ) may increase intestinal absorption increasing blood levels and the risk of bleeding. Conversely P-gp inducers (rifampicin, carbamazepine and phenytoine) may decrease plasma concentrations of dabigatran. Apixaban and rivaroxaban are substrates of CYP3A4 / 5 and P-gp transporters. Concomitant administration of inhibitors of these systems such azole antifungine, HIV protease inhibitors may lead to increase of anticoagulation. By converse inducers of these systems decrease plasma levels of apixaban and rivaroxaban. No significant increase of bleeding rate has been demonstrated with concomitant use of aspirin or NSAID in patients treated with dabigatran both at 150 and 220 mg doses.

![Figure 1. Mechanism of action of new anticoagulants](image)
Anti thrombin agent

Dabigatran etexilate is a pro-drug quickly converted in dabigatran by cytochrome P-450–independent esterases. Anticoagulant effect of dabigatran is related to a powerful reversible direct competitive inhibitor of both free and clot included thrombin. Onset of action is prompt, it has a consistent anticoagulant effect. The half-life ranges between 12 and 17 h [40] but since kidney excretion accounts for 85% elimination, half-life does increase significantly also in patients with moderate renal failure (creatinine clearance < 50 ml/min).

Two large randomized trials compared dabigatran etexilate 75 mg or 110 mg within 1 to 4 hours of surgery followed by 150 or 220 mg daily thereafter to enoxaparin 40 mg on the day prior to surgery and daily thereafter in patients undergoing elective major orthopedic surgery. In the RE-MODEL trial were included 2101 patients undergoing knee replacement [41] and duration of treatment was 6 to 10 days. Exclusion criteria were haemorrhagic diathesis and severe liver or kidney failure. Administration of COX-2 inhibitors and aspirin were allowed during treatment period. Dabigatran etexilate at 150 and 220 mg was statistically non-inferior to enoxaparin in the combined end point of symptomatic or asymptomatic (diagnosed by venography at the end of treatment period) DVT, non fatal pulmonary embolism and all cause mortality. In RE-MODEL study symptomatic VTE occurred in 2.6% and 3.8% at dabigatran dosage of 220 mg and respectively 150 mg in comparison to 3.5% of enoxaparin. In REMOBILIZE study [42] dabigatran (150 or 220 mg) was compared to enoxaparin (30 mg b.i.d.) for prevention of VTE after TKR. At variance with other studies enoxaparin was superior to dabigatran. However better results obtained with enoxaparin may be explained by the higher doses employed and by a later start of treatment with dabigatran (12 hours after surgery).

A decreased dose (150 mg daily) must be prescribed in patients aged > 75 years or with impaired renal function (creatinine clearance between 30 and 60 ml/min) or treated with P-glycoprotein inhibitors (verapamil, amiodarone or quinidine).

Activated X factor inhibitors

Rivaroxaban is an active direct inhibitor of coagulation Factor Xa. Maximal inhibition of factor Xa occurs three hours after a dose. Although effects lasts 8–12 hours factor Xa activity does not return to normal within 24 hours so once-daily dosing is advised.

The effects of rivaroxaban in orthopedic surgery have been examined RECORD studies. In this group of trials primary composite efficacy endpoint was any DVT, nonfatal PE, or death from any cause. The RECORD 3 (2531 patients) study [45] showed that rivaroxaban started postoperatively and administered for 10–14 days was significantly more effective than enoxaparin 40 mg/day started preoperatively in patients undergoing TKA. The absolute risk reduction of the primary endpoint was 9.2% at two weeks in RECORD 3. Safety profiles were similar between the two drugs.

Rivaroxaban was compared with enoxaparin 30 mg b.i.d, both started postoperatively and continued for 10–14 days in patients undergoing TKR in RECORD 4 trial [44]. Incidence of primary efficacy outcome at day 17 after surgery were respectively 6.9% and 10.1% in patients randomized to rivaroxaban or enoxaparin. After introduction of NICE guidelines administration of rivaroxaban was associated, in comparison to an historical control group, with a significant decrease of symptomatic confirmed pulmonary embolism (2/266 vs 24/596, p=0.0084) but at the expense of a significant increase of complications at surgical site [45].

The significant decrease of total DVT and all cause mortality (HR 0.46, 95% CI 0.39-0.54), was obtained however at the expense of a significantly higher risk of significant bleeding and a trend towards an higher risk of major bleeding (RR 1.25, 95% CI 1.05-1.49). A recent retrospective study compared 369 patients receiving simultaneous combined mechanical and pharmacological thromboprophylactic modality and 385 patients receiving sequential combined modality with early-mechanical compression treatment followed by rivaroxaban 2 days later after primary total knee arthroplasty surgery [46]. The incidence rates of proximal and distal DVT tested on the 2nd postoperative day, had no significant difference in the sequential group (4.16%) compared with those (3.25%) in the simultaneous group. No difference was found in the incidence risk of proximal and distal DVT on the 5th week between groups. No pulmonary embolism occurred. The mean volume of wound drainage in the sequential group was 345 mL which was 98 mL lesser than in the simultaneous group (p < 0.001). The use of the mechanical compression method alone during the early-postoperative 48 hours, then followed by rivaroxaban then until the 5th week had the same antithrombotic effects and reduced the postoperative wound drainage volume as compared with simultaneous combined modalities.

Apixaban

Apixaban is a reversible inhibitor of activated X factor. Apixaban half life is close to 12 hours. The drug is cleared through hepatic and renal route, the last contributing for 27%. Daily dosage is 2.5 mg b.i.d. The effects of apixaban in major orthopedic surgery were examined by the ADVANCE trials. In ADVANCE 1 apixaban 2.5 mg b.i.d. was compared to enoxaparin (30 mg every 12 hours started after surgery) in 3195 patients undergoing TKA [47]. Treatment were administered for 10-14 days. Primary end point (symptomatic or asymptomatic DVT, non fatal EP and overall mortality) was reached in 9% of apixaban group in comparison to 8.8% in enoxaparin group. Bleeding was less frequent with apixaban than with enoxaparin - 0.7% vs. 1.4% (p = 0.053).

ADVANCE 2 trial (3057 patients) compared apixaban with enoxaparin in patients undergoing TKA. Primary efficacy
outcome incidence was 15.1\% in the apixaban group and 24.4\% in the enoxaparin group (relative risk 0.62, 95\% CI 0.51–0.74) [48]. In apixaban group proximal DVT, symptomatic nonfatal PE, and VTE-related death occurred in 1.1\% of patients vs. 2.2\% of those given enoxaparin (relative risk 0.50, 95\% CI 0.26–0.97). The incidence of bleeding was lower with apixaban in comparison to enoxaparin (P = 0.09).

A metaanalysis of the ADVANCE studies showed that apixaban (2.5 mg b.i.d.) compared to enoxaparin significantly decreased total DVT and all cause mortality (HR 0.46, 95\% CI 0.39–0.54). However the risk of symptomatic VTE was not significantly different (RR 0.81, 95\%CI 0.41–1.66, p= 0.57). Otherwise the administration enoxaparin was associated with an higher incidence of bleeding requiring reoperation. The reported lower risk of major bleeding and significant bleeding reported for apixaban is limited by the fact that drop in haemoglobin was not calculated in comparison to baseline values but on immediate postoperative haemoglobin, thus effective bleeding effects may be underestimated.

**Edoxaban**

Limited experience has been provided with edoxaban in elective orthopaedic surgery. STARS E-3 compared edoxaban (30 mg once daily started 6-24 hours after surgery) with enoxaparin 20 mg b.i.d, in 746 patients undergoing TKA [49]. In STARS E-3 edoxaban decreased DVT in comparison to enoxaparin (2.4 vs 6.9\%). Bleeding rate did not differ between the two drugs. Recently were published data about 300 patients undergoing TKA treated with edoxaban 15 mg , enoxaparin 20 mg b.i.d and fondaparinux 1.5 mg once a day for a period of 14 days [50]. Edoxaban resulted more efficient in comparison to the other two prophylaxis regimen. A randomized study evaluated the effectiveness of a A-V Impulse System foot pump associated with edoxaban in the prophylaxis of DVT after TKA [51]. The authors did not find any significant difference in primary outcomes (any DVT as detected by bilateral ultrasonography up to postoperative day 10 and pulmonary embolism PE up to postoperative day 28) and safety outcomes (bleeding and death of any cause up to postoperative day 28).

**Management of bleeding**

Specific antidotes for NAO are in development and idarucizumab, a drug-specific antidote targeted to reverse the direct thrombin inhibitor dabigatran has been demonstrated efficacy and safe in animal models [52] and in healthy volunteers [53]. Idarucizumab administration is associated with immediate, complete, and sustained reversal of dabigatran-induced anticoagulation in healthy men, and was well tolerated with no unexpected or clinically relevant safety concerns. Andexanet alfa is a class-specific antidote targeted to reverse the oral direct factor Xa inhibitors as well as the indirect inhibitor, enoxaparin. Ciraparantag is a universal antidote targeted to reverse the direct thrombin and factor Xa inhibitors as well as the indirect inhibitor, enoxaparin. At present, waiting clinical introduction of drug antidotes, the management of bleeding should be guided by location and severity of haemorrhage. Drug discontinuation and local compression, when possible, should be immediately adopted to stop bleeding. In case of minor bleeding (haematuria or epistaxis) resuming drug after resolution of bleeding may be sufficient. In patients with major bleeding maintaining diuresis, administration of charcoal, transfusion of red blood cells and fresh frozen plasma may be used. Limited evidence exist pro that thrombomodulin complex concentrates, activated prothrombin complex concentrates (FEIBA) and rVIIa factor, may contribute to reversing the anticoagulant effects of Xa factor inhibitors in patients with severe hemorrhagic complications.

**TIMING OF DRUG ADMINISTRATION FOR THROMBOPROPHYLAXIS**

Timing of the first dose of anticoagulants in patients undergoing TKR has not still clearly defined. In TKR perioperative thromboprophylaxis, 2 hours before till to 6 hours after surgery, has been associated with increase of bleeding risk [54]. Late treatment, beyond 12 hours, otherwise seems to be associated with an higher VTE risk.

**COMPARISON OF DIFFERENT METHODS**

Few studies compared the efficacy and safety of different antithrombotic methods and/or agents on venous thromboembolism (VTE) after unilateral total knee arthroplasty. A recent study from China showed that aspirin, low-molecular-weight heparin, and rivaroxaban could effectively reduce the incidence of VTE after total knee arthroplasty, and their efficacy was similar [55]. Rivaroxaban however had a higher incidence of bleeding complication suggesting again some concerns on its safety.

Bern et al [56] randomized 355 patients in three groups: A: variable dose warfarin (first dose on the night preceding surgery with subsequent target INR 2.0–2.5), B: 2.5 mg fondaparinux daily starting 6-18 h postoperatively, or C: fixed 1.0 mg dose warfarin daily starting 7 days preoperatively. All treatments continued until bilateral leg venous ultrasound day 28±2 or earlier upon a VTE event. Two patients in Arm C had asymptomatic distal DVT. One major bleed occurred in Arm B and one in Arm C (ischemic colitis). Fixed low dose warfarin started preoperatively was equivalent to two other standards of care under study (95\% CI: 0.0428, 0.0067 for both) vs VTE prophylaxis.

A metaanalysis of three studies involving 11,659 patients, the risk of symptomatic DVT (pooled OR 0.38, 95\% CI 0.16-0.90, , p=0.03) and bleeding (pooled OR 0.87, 95\% CI 0.77-0.99, p=0.03) was less in apixaban group compared to the enoxaparin group [57]. However, it was interesting to note that on subgroup analysis, the risk of PE was higher with apixaban when used for thromboprophylaxis in knee replacement surgery (pooled OR 2.58, 95\% CI 1.10-6.04, p=0.03).
VTE PROPHYLAXIS IN TKR IN HAEMOPHILIA

A debated point is the need of a VTE prophylaxis in patients affected by Haemophilia undergoing orthopaedic surgery [58].

Haemophilia is a X-linked disease characterized by an inherited lack of coagulative factors as factor VIII or IX associated with a high risk of haemorrhages. Due to the recent improvements in the prevention of bleedings by primary or secondary haematologic prophylaxis, the most frequent complication of Haemophilia is the haemophilic arthropathy in specific joints, called “target joints” [59]. The most involved joint is the knee, and often the quality of life is poor due to the severity of symptoms and functional impairment: conservative treatments and minimally invasive procedures are rarely useful in such cases [60-61]. In the recent decades a high increase of the TKA performed in such patients has been recorded, given the fact that major surgery is the only procedure that may induce adequate pain relief and functional recovery [62-63].

VTE is a recognized complication after joint replacement surgery, and a prophylaxis is routinely used in patients without bleeding disorders. However, in haemophilic patients, the pharmacologic VTE prophylaxis is highly variable and controversial because of the inherent bleeding risk.

A recent review regarding 71 hip or knee replacements in 42 consecutive patients affected by Haemophilia A or B was compared to the literature in order to ascertain the incidence of VTE after THA and TKA in the haemophilic population [58]. In all patients compression stockings were applied up to 6 weeks after surgery; additionally, in 6 cases (10.5%) sequential intermittent compression devices were used, and 2 (2.8%) received a postoperative low-molecular-weight heparin (LMWH) regimen. One patient (1.4%) receiving LMWH had a symptomatic VTE ten days after the hip replacement for a traumatic proximal femur fracture. None of the remainders (70 patients) reported any symptomatic VTE within 3 months after surgery. An estimated incidence of symptomatic VTE of 0.5% was assessed from the analysis of pooled data from several series of haemophilic patients undergoing a joint replacement. The conclusion was that in patients with Haemophilia, joint replacements can be safely performed without a routinary VTE prophylaxis, and without an increasing risk of thromboembolic events [58].

Another retrospective study on 38 cases of TKA in 33 haemophilic patients was reported to evaluate any case of DVT pre- and postoperatively by US. No patients performed a DVT prophylaxis. Also D-dimer and fibrinogen degradation product were measured before and after surgery. No case of DVT was found in either pre- or postoperative examinations [64].

This approach has been safely adopted in many series and not only for knee surgery, but also for other major procedures, as Total Hip Arthroplasty (THA) [65].

Peculiar conditions in which a VTE prophylaxis may be not indicated is the presence of inhibitors, antibodies against recombinant factors developed by some patients, able to limit the efficacy of clotting factors [66-67]. In such patients, the high risk of bleeding is more consistent than the risk of VTE. However, the high doses of bypassing agents (recombinant factor VII) used to activate the coagulative cascade may induce a variable risk requiring a strict and multidisciplinary daily evaluation in dedicated centres by tailored administration protocols.

To reduce the risk of VTE in inhibitor patients undergoing major orthopaedic surgery, the use of non-pharmacologic measures, such as standard compression stockings, mechanical prophylaxis, and early functional recovery seem to be appropriate.

CONCLUSIONS

The synthesis of recommendation for VTE prophylaxis in knee replacement has been provided by guidelines produced in the recent past years by international scientific societies [66-67]. The American college of Chest Physicians antithrombotic and prevention of thrombosis guidelines [69] suggests that in patients undergoing TKA the administration of LMWH, fondaparinux, apixaban, dabigatran, rivaroxaban, LDH, adjusted dose warfarin of finally aspirin has grade 1 B recommendation, while intermittent compression devices (ICPD) has a grade 1c recommendation. The use of LMWH (with or without ICPD) is preferred to other drugs with a grade 2 B recommendation. LMWH should be preferred to other agents in patients at high risk of bleeding. Since a non-adherence rate between 13 and 27% has been demonstrated in patients treated with LMWH or fondaparinux, direct oral anticoagulants should be considered in patients with expected poor adherence to parenteral route of administration. The treatments should be started either 12 or more hours before surgery or 12 hours or more after. Combination of pharmacological prophylaxis and ICPD for at least 18 h/day has a grade 2 C recommendation. ICPD alone may be considered for patients at high risk of bleeding. IVC filters are not recommended for prophylaxis of thromboembolism in patients undergoing major orthopaedic surgery, even in the case of high risk of bleeding or contraindication to mechanical prophylaxis. American Academy of Orthopedic Surgeons (AAOS) guidelines for the prevention of venous thromboembolism (VTE) in patients undergoing hip or knee surgery [67] evaluate the efficacy of various agents in preventing pulmonary embolus rejecting asymptomatic DVT as end point. The AAOS guidelines accept the use of aspirin combined with mechanical compression devices, and low-dose warfarin therapy. These guidelines in synthesis express a major concern with the risks associated with anticoagulation therapy. However as expressed by ACCP guidelines randomized data clearly demonstrated a decrease of both DVT and PE by thromboprophylaxis. Moreover the AAOS recommended routine preoperative assessment for risk of PE and bleeding and stratified their recommendations for thromboprophylaxis according to whether patients are “standard risk” or “high risk” for PE and bleeding. They did
not discuss the issue of dose adjustments in the elderly or those with renal dysfunction, which are the most important indicators of high bleeding risk.

Low molecular weight heparin at European dosage (4000 IU) started before surgery is the usual regimen adopted in elective knee arthroplasty in our clinical practice while direct oral anticoagulants are prescribed in selected cases with expected poor compliance to parental agents.

CONFLICT OF INTEREST
The authors declare that they have no conflict of interest

LIST OF ABBREVIATIONS:

DVT = deep venous thrombosis
THA = total hip arthroplasty
TKA = total knee arthroplasty
LMWH = low molecular weight heparin
UI = un fractionated heparin
PTS = post thrombotic syndrome

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