Introduction

After neurosurgical procedures with long general anesthesia, deep venous thrombosis (DVT) and pulmonary embolism (PE) are major causes of postoperative morbidity and mortality. In untreated control groups of randomized studies after craniotomies, DVT rates exceed 25%.\(^1\) In such patients, clinically evident DVT occurs in 2% to 4%.\(^2\) PE rates vary between 0.8% and 2%.\(^3,4\) The efficiency, cost-effectiveness, and safety of prophylactically administered anticoagulative drugs after general surgical and neurosurgical procedures have been demonstrated in clinical studies.\(^5-10\) Furthermore, the...
The additive use of mechanical devices such as elastic graduated compression stockings or pneumatic compression boots has been shown to reduce the incidence of DVT after surgical procedures. However, a standardized protocol in the perioperative prevention of DVT for neurosurgical procedures is still missing. For one, this is because of the fear of secondary hemorrhage, which is supposed to be increased when applying anticoagulative drugs early in the postoperative course. Furthermore, the available literature does not yield a consensus concerning the management of postoperative DVT or PE with therapeutic anticoagulation after craniotomies.

### Methods

A consecutive series of patients undergoing craniotomies between 2006 and 2011 with postoperative PE or DVT was retrospectively analyzed regarding demographic attributes, diagnosis, method of perioperative thrombosis prevention, and interval between surgery and PE or DVT. PE and DVT were detected by elevated d-dimer levels, ultrasonography, and lung computed tomography (CT). In addition, the administered anticoagulative medication and its dosage were examined as well as patients’ weight and kidney and liver functions. Secondary hemorrhage was excluded either by uneventful course or by cranial CT. Routine postoperative CT was not performed. The decision for postoperative CT was made individually and after consultation with the surgeon. The surgeon decided on the dosage of the applied anticoagulative drugs depending on the assumed risk for a secondary hemorrhage in each individual case. Subtherapeutic anticoagulation was defined as a dosage higher than prophylactic in contrast to therapeutic heparinization as the standard dosage for treatment of PE or DVT in a nonsurgical setting.

### Results

A consecutive series of 42 patients was retrospectively analyzed. Indications for surgery were meningiomas (n = 17), vestibular schwannomas (n = 8), aneurysms (n = 5), gliomas (n = 4), subdural hematomas (n = 3), ependymoma (n = 1), hemangioblastoma (n = 1), brain metastases (n = 1), prolac- tinoma (n = 1), and intracerebral hemorrhage (n = 1). A total of 26 patients were female, and 16 were male. Patients’ age ranged from 17 to 80 years (mean, 55 years). None of the patients had a severe impairment of kidney or liver function.

Prevention of postoperative thromboembolism was performed with elastic graduated compression stockings and enoxaparin, certoparin, or unfractionated heparin (UFH). Low-molecular-weight heparin (LMWH) for standard perioperative prophylaxis was given starting on the first postoperative day. In some cases of, for example, intracerebral or subdural hematoma, the start of LMWH administration was determined individually. None of the patients received anticoagulative drugs on the day of surgery. Prophylactic medication was started on the first postoperative day in 33 patients, on the second postoperative day in two patients, on the third postoperative day in three patients, on the tenth postoperative day in one patient, on the eleventh postoperative day in two patients, and on the twelfth postoperative day in one patient.

Postoperative PE or DVT was observed in 41 patients between the first and the 28th postoperative day (median, fifth postoperative day) (Fig. 1). Seven patients suffered from PE and DVT, one patient only from PE, and 34 patients exclusively from DVT. In one patient (number 9) a DVT 12 days before surgery was diagnosed, resulting in higher pre- and postoperative dosages of enoxaparin (Fig. 1).

During the postoperative course, 19 patients developed symptoms of DVT or PE (dyspnea, pain or swelling in the lower leg), and 23 did not have any symptoms. In these 23 patients, pre- and postoperative d-dimer levels were measured according to a protocol of a separate study investigating postoperative d-dimer values. In case of distinct postoperative elevation of d-dimer levels, deep vein ultrasonography or lung CT scans were performed to verify PE or DVT.

After the diagnosis of postoperative PE or DVT, the dosage of enoxaparin or partial thromboplastin time (PTT)-monitored UFH was increased in 39 patients. Prophylactic anticoagulation was continued in the three remaining patients because of the small extent of DVT. Therapeutic heparinization was applied to 30 of the 42 patients.

Assuming that the risk for a secondary hemorrhage is higher in the early postoperative course, the first week after surgery was analyzed in detail. During the first 7 days after surgery (starting on postoperative day 2 to 7), 20 patients were treated with higher than prophylactic dosages. Seven of them received anticoagulative drugs in dosages of standard therapeutic heparinization, and 13 patients received subtherapeutic dosages between prophylactic and therapeutic heparinization. For anticoagulation, LMWH (enoxaparin subcutaneously 2 – 30 mg to 2 – 70 mg) or UFH (target PTT, 60–80 sec) delivered through a syringe pump were used (Fig. 2). Fig. 2 illustrates the management of anticoagulation in relation to the interval between craniotomy and start of therapeutic/subtherapeutic heparinization. Lung CT scans revealed extended bilateral PE in 5 of these 20 patients. The remaining 15 patients exclusively had DVT.

None of the 42 patients had a secondary hemorrhage after LMWH or heparin administration. Secondary hemorrhage
Discussion

The general risk for secondary hemorrhage after intracranial procedures was reported at 1.5% to 2.5%. Administration of heparin 12 hours after intracranial surgery is relatively safe. In contrast, administration of enoxaparin in prophylactic dosages during induction of anesthesia was associated with an increased risk for secondary hemorrhage after intracranial tumor surgery. Consequently, the interval between surgery and the start of anticoagulative medication seems to be more important than its dosage regarding the risk for secondary hemorrhage. In a prospective, randomized, double-blind study, there was no significant difference concerning hemorrhage complications between patients treated with subcutaneous heparin and those who were not. In contrast, a higher but not statistically significant hemorrhage complication rate (2.5% vs. 0.8%) was reported for the prophylactic use of LMWH. A randomized pilot study showed no differences in postoperative hemorrhage, DVT, or PE between the group treated with subcutaneous heparin and the group with LMWH medication. Prophylactically administered LMWH starting within 24 hours after intracranial procedures reveals a risk reduction of 28.9% for DVT and of 40.2% for PE. Enoxaparin with compression stockings was more effective than compression stockings alone for prevention of DVT in elective neurosurgical procedures. Major hemorrhage complication rates of 3% were observed each in the enoxaparin and the placebo group. Cerrato et al reported a risk reduction from 34% to 6% for DVT by prophylactically administered heparin and no significantly increased rate of postoperative hematomas. A randomized, prospective, double-blind clinical trial revealed the efficacy and safety of enoxaparin and UFH for prophylaxis of venous thromboembolism after brain tumor surgery. However, heparin-induced thrombocytopenia is a potential adverse drug effect of heparin medication with a reported frequency of 0.2% to 5%. After LMWH administration, the risk is five- to 10-fold lower compared with UFH.

A recently published study reviewed the current practice of perioperative prevention of DVT in German neurosurgical departments. There was no homogenous practice in the administration of heparin (UFH, LMWH, or both) and even compression stockings. In general, the risk for secondary hemorrhage under heparin was estimated higher after cranial versus spinal interventions. Heparin application after craniotomy starting during the first 5 days after surgery is assumed to be associated with a high or very high risk for relevant secondary hemorrhage by up to 20% of German neurosurgical departments.

Although a retrospective study cannot provide sufficient evidence of higher than prophylactic dosages of anticoagulative drugs being a safe treatment of PE or DVT after craniotomies, our results suggest that the rate of secondary hemorrhages under these circumstances may be overestimated. As in studies dealing with prophylactically administered low-dose heparin, no significant postoperative hemorrhages were observed in the present study analyzing therapeutic anticoagulation after craniotomies. Taking into consideration the potential risk for postoperative secondary hemorrhages, the risk-to-benefit ratio for using anticoagulative drugs in therapeutic dosages in patients with PE or DVT after craniotomy may be still favorable for the patient.

Our review of the literature and MEDLINE research revealed no further studies concerning therapeutic anticoagulation in cases of PE or DVT after craniotomies. In general, neurosurgeons hesitate to administer higher than prophylactic dosages of anticoagulative drugs in the early postoperative course after craniotomies. The reasons might be fear of secondary hemorrhages, the lack of clinical studies and accepted protocols in the postoperative management of PE and DVT after craniotomies and medicolegal considerations because LMWH and heparin are not approved after neurosurgical procedures.

The presented study is the first published clinical series of patients with LE or DVT after craniotomy receiving LMWH. The old data of Swann et al about management of LE or DVT lacks comparability because of the administered anticoagulative substances and the inclusion of patients with spinal disorders. The patient number in this study with DVT or LE after craniotomy was 10, and the delay between surgery and beginning of anticoagulative medication was 8 to 54 days (mean, 26 days). Rebleeding was only observed in a patient
without surgery. Ruff et al. included more patients but used the same anticoagulative regimen as Swann et al but no LMWH. The prospective study of Gerlach et al. reporting on 2823 patients undergoing major and minor intracranial procedures dealt with the prophylaxis of thromboembolic events. The safety of therapeutic heparin administration in patients diagnosed with DVT or PE in the course was not addressed, as in the study of Goldhaber et al. and the review article of Epstein.

In 14 patients in the present study, no postoperative CT scans were performed. These patients might have developed secondary hemorrhages. Yet without clinical signs, the radiologic finding of a small secondary hemorrhage would not have had any effects on treatment. Additionally, although limited by the variety of applied substances and their dosages, our data suggest no differences between the used anticoagulative drugs concerning the risk for rebleeding. Consequently, systematic studies are needed to estimate the risk for secondary hemorrhages under therapeutic anticoagulation after craniotomy.

**Conclusion**

The documented differences in the used anticoagulative substance, the start of medication, and the dosage reflect the need for a standardized protocol concerning the treatment of postoperative PE or DVT after craniotomy. Considering there was no secondary hemorrhage in this study and that PE and DVT are potentially life-threatening complications, it may be justified to be more aggressive in the application of anticoagulative drugs after craniotomy. Our observations do not support the assumption of a very high risk for rebleeding with application of heparin in the early days after craniotomy.

**Acknowledgments**

We thank Christin Zöller for her secretarial contribution and assistance in literature review.

**Conflict of Interest**

None

**References**


24 Martel N, Lee J, Wells PS. Risk for heparin-induced thrombocytope-
nia with unfractionated and low-molecular-weight heparin throm-
25 Swann KW, Black PM, Baker MF. Management of symptomatic
depth venous thrombosis and pulmonary embolism on a neuro-
26 Ruff RL, Posner JB. Incidence and treatment of peripheral venous
postoperative hemorrhage after intracranial surgery after early
nadroparin administration: results of a prospective study. Neuro-
28 Epstein NE. A review of the risks and benefits of differing prophyl-
axis regimens for the treatment of deep venous thrombosis and
pulmonary embolism in neurosurgery. Surg Neurol 2005;64(4):
295–301, discussion 302