Prevention of venous thromboembolism in patients undergoing bariatric surgery

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Abstract: Bariatric surgical procedures are now a common method of obesity treatment with established effectiveness. Venous thromboembolism (VTE) events, which include deep vein thrombosis and pulmonary embolism, are an important source of postoperative morbidity and mortality among bariatric surgery patients. Due to an understanding of the frequency and seriousness of these complications, bariatric surgery patients typically receive some method of VTE prophylaxis with lower extremity compression, pharmacologic prophylaxis, or both. However, the optimal approach in these patients is unclear, with multiple open questions. In particular, strategies of adjusted-dose heparins, postdischarge anticoagulant prophylaxis, and the role of vena cava filters have been evaluated, but only to a limited extent. In contrast to other types of operations, the literature regarding VTE prophylaxis in bariatric surgery is notable for a dearth of prospective, randomized clinical trials, and current professional guidelines reflect the uncertainties in this literature. Herein, we summarize the available evidence after systematic review of the literature regarding approaches to VTE prevention in bariatric surgery. Identification of risk factors for VTE in the bariatric surgery population, analysis of the effectiveness of methods used for prophylaxis, and an overview of published guidelines are presented.

Keywords: bariatric surgery, venous thromboembolism, prophylaxis, vena cava filter, heparin

Introduction

Surgical approaches to weight loss, bariatric surgeries, are commonly performed procedures for morbidly obese individuals; the estimated number of bariatric procedures in the USA alone was close to 180,000 in 2013. Bariatric surgery is effective in achieving weight loss and improving obesity-related complications. However, there are also potential risks or complications, among them venous thromboembolism (VTE). Reported rates of VTE, including deep vein thrombosis (DVT) and pulmonary embolism (PE), following bariatric surgery are 0.3%–2.2%, with rates of PE being approximately 1%, despite application of methods to prevent these complications. PE is a frequent cause of postoperative mortality in the bariatric surgery population and is a common finding at autopsy.

Various strategies have been used to prevent VTE in patients undergoing bariatric surgery, including pharmacologic and mechanical approaches. However, the optimal approach remains unclear. The objective of this review is to discuss and evaluate the existing literature regarding prevention of VTE in bariatric surgery patients. This review includes identification of risk factors for VTE in the bariatric surgery population,
analysis of the effectiveness of methods used for prophylaxis, and an overview of published guidelines.

Methods
Search strategy
A systematic literature search was performed in MEDLINE (1946–January 12, 2015), EMBASE (*1947–January 12, 2015), Cochrane Database for Systematic Reviews, and Clinicaltrials.gov and is depicted in Table 1. The search was limited to English language studies. Search terms for bariatric surgery included both generic and specific terms for various bariatric procedures; broad terms for venous thrombosis and methods of prophylaxis were used.

Study selection and data abstraction
Inclusion and exclusion criteria to identify studies for this review are shown in Table 2. The primary aim of these criteria was to identify studies which report outcomes from two or more treatment groups or include multivariate analysis to control for VTE risk factors. Abstracts not published as peer reviewed articles were not included.

Following the primary literature search, two reviewers (MB and PD) independently screened all articles to ensure satisfaction of inclusion/exclusion criteria. Reference lists of included articles were reviewed to identify additional publications of interest. Information on general study characteristics (study design, number of participants, study period, and follow-up), study participants (age, sex, body mass index [BMI]), bariatric procedure type, methods of prophylaxis, and outcome measures was collected.

Results of study selection
A summary of the selected studies is presented in Tables 3 and 4 (Table 3: mechanical and pharmacologic prophylaxis studies; Table 4: vena cava filter [VCF] studies). Regarding study designs used, only two were prospective, randomized trials; all others were either retrospective or prospective cohort studies. Most studies were single center, although there were large database driven studies included. Sample sizes were relatively small with respect to numbers needed to detect differences in the primary outcome metrics (VTE). In some cohort studies, different treatment groups were recruited consecutively over several years.

Modes of prophylaxis reported include lower extremity compression (LEC), anticoagulation therapy including subcutaneous (SC) low-molecular-weight heparins (LMWHs) and unfractionated heparin (UFH), and VCFs. Multiple studies addressed the question of adjusted-dose heparin. A few of the studies of pharmacologic prophylaxis included patients who also received a VCF, though the proportions with VCF were small.

Roux-en-Y gastric bypass (RYGB), either laparoscopic or open, was the only procedure type in multiple studies. The proportion of open versus laparoscopic of any procedure type was not always reported. A few studies did not describe patient age, proportion of male participants, or BMI of study subjects. In some cases, patients with a history of VTE were excluded, and in some cases, the frequency of patients with a VTE history was not reported. In many studies, comparisons of the frequencies of patient VTE risk factors or of procedure type are not reported or tested.
All studies except one included in this review report VTE outcomes. In general, VTE events were identified based on testing directed to symptomatic patients or clinical suspicion of disease, although two studies performed imaging to detect asymptomatic DVT. PE was diagnosed through evaluation of symptoms. Bleeding events are reported in studies in which the primary aim was evaluation of pharmacologic prophylaxis. However, there is no standard definition of bleeding severity across these reports, making comparisons challenging. The majority of studies report postoperative mortality, but do not always distinguish PE-related death from other causes. Filter-related complications are not consistently reported across the studies of VCF as prophylaxis. The postoperative outcome ascertainment period for most studies was between 30 and 90 days.

**Risk factors for VTE among bariatric surgery patients**

Risk factors for postoperative VTE have been identified for surgical patients and have been incorporated into validated risk assessment tools. An examination of VTE risk factors specific to the bariatric surgery population is warranted; this understanding provides insight into the bariatric surgery prophylaxis literature and could refine future prevention strategies. Postoperative VTE risk factors can be categorized as patient related or procedure related; identified risk factors from at least one published report are presented in Table 5.

**Patient-related risk factors for VTE**

Among patient-related characteristics, multiple studies have found that the male sex is associated with an elevated risk of VTE among bariatric surgery patients. Preoperative patient weight and BMI have also been associated with an increased risk of VTE events. For example, Finks et al demonstrated that every 10 unit increment in BMI was associated with a 37% increase in VTE risk (relative risk [RR], 1.37; confidence interval [CI], 1.06–1.75). Several studies have also identified increasing age as a risk factor for postoperative VTE. Patient smoking status has been identified as a potential VTE risk factor in two reports. Several of the previously referenced studies have identified a prior history of VTE as a predictor of postoperative VTE in the bariatric surgery population. For example, Finks et al demonstrated that a prior history of VTE was associated with four times the risk of postoperative VTE (odds ratio [OR], 4.15; CI, 2.42–7.08). Studies have also evaluated the presence of possible markers of hypercoagulability among bariatric surgery patients but have not assessed an association with clinical VTE.

**Procedure-related risks factors for VTE**

Procedure-related factors for VTE after bariatric surgery include operative time, procedure type, postoperative complications, and whether the procedure is open or laparoscopic. In the bariatric surgery literature, studies have indicated an increased risk of VTE with open compared with laparoscopic procedures. Regarding duration of surgery, Finks et al reported an 86% increased risk of VTE with operative time >3 hours, (RR, 1.86; CI, 1.07–3.26). Chan et al also identified operative time >3 hours as an independent predictor of postoperative VTE and found that preoperative BMI was an independent predictor of operative time. In the analysis by Jamal et al, revision surgeries were associated with elevated VTE risk. The report by Gonzalez et al found that postoperative anastomotic leak after RYGB was also associated with increased VTE risk. In a comparison of different types of bariatric procedures, Finks et al found higher VTE risk with sleeve gastrectomy, laparoscopic gastric bypass, open RYGB, and duodenal switch surgery when compared to adjustable gastric band procedures. Masoomi et al found that gastric bypass procedures carry higher VTE risk when compared to other types of bariatric procedures.

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**Table 2 Inclusion and exclusion criteria for identified publications**

| Inclusion criteria                                                                 | Exclusion criteria                                                                 |
|-----------------------------------------------------------------------------------|------------------------------------------------------------------------------------|
| • Human subjects                                                                  | • Animal studies                                                                   |
| • Patients undergoing bariatric surgery                                            | • Study subjects <18 years of age                                                  |
| • Study designs: randomized trials, or cohort studies comparing two or more groups | • Patients undergoing contouring and plastic surgery following bariatric surgery    |
| • Studies comparing VTE prophylaxis strategies: lower extremity compression, pharmacologic prophylaxis, or vena cava filters | • Abstract only (no peer-reviewed published article)                                |
| • Studies reporting postoperative clinical outcomes: venous thromboembolism, bleeding complications, or mortality | • Observational studies with no control or comparison group or study data presented without multivariate analysis |

**Abbreviation:** VTE, venous thromboembolism.
| References               | Design                  | Patients (N) | Method(s) of prophylaxis analyzed | Dosing schedule for anticoagulant | Duration of prophylaxis (days) | VTE events (%) | Bleeding events (%) | Mortality (%) |
|--------------------------|-------------------------|--------------|-----------------------------------|-----------------------------------|------------------------------|----------------|---------------------|---------------|
| Gagner et al (2012)      | Prospective cohort      | 4,020        | SCD and pharmacologic prophylaxis (UFH or LMWH) | Not specified                      | NR                           | 0.47            | NR                  | 0.35           |
| Frantzides et al (2012)  | Prospective cohort      | 396          | SCD                               | NA                                | NR                           | 0.25            | NR                  | 0.25           |
|                          |                         | 435          | SCD and LMWH                       | Enoxaparin 40 mg q12h              | NR                           | PE 1.1*         | 4.8*                | 0.12 (not VTE related) |
|                          |                         | 1,257        | SCD; LMWH to high risk only (personal or family history of hypercoagulable state or family history of VTE) | Enoxaparin 40 mg q12h              | NR                           | PE 0*           | 0.4*                | 0             |
| Birkmeyer et al (2012)   | Prospective cohort      | 4,402        | Pre- and postoperative UFH         | NR                                | NR                           | 0.68*           | 0.46                | NR            |
|                          |                         | 4,482        | Preoperative UFH/postoperative LMWH | NR                                | NR                           | 0.29*           | 0.6                 | NR            |
|                          |                         | 15,891       | Pre- and postoperative LMWH        | NR                                | NR                           | 0.25            | 0.38                | NR            |
| Kothari et al (2007)     | Prospective cohort      | 238          | LMWH                              | Enoxaparin 40 mg q12h              | Preoperative to discharge     | 0               | 5.9*                | 0             |
|                          |                         | 238          | UFH                               | 5,000 U q8h                        | Preoperative to discharge     | 0.42            | 1.3*                | 0             |
| Imberti et al (2014)     | Prospective randomized trial | 131         | Pre- and postoperative LMWH       | Parnaparin 4,250 U q24h            | 14.1±2.4 (mean ± SD)          | PE 0.76         | 6.1                 | 0             |
|                          |                         | 119          | Pre- and postoperative LMWH       | Parnaparin 6,400 U q24h            | 14±2.5 (mean ± SD)            | PE 0            | DVT 0.84            | 5.0           |
| Kalfarentzos et al (2001)| Randomized controlled trial | 30          | Pre- and postoperative LMWH       | Nadroparin 5,700 q24h              | Preoperative to discharge     | 0               | 0                   | 0             |
|                          |                         | 30           | Pre- and postoperative LMWH       | Nadroparin 9,500 q24h              | Preoperative to discharge     | 0               | 6.7                 | 0             |
| Scholten et al (2002)    | Prospective cohort      | 92           | Pre- and postoperative LMWH       | Enoxaparin 30 mg q12h              | NR                           | 5.4             | 1.1                 | NR            |
| Hamad and Choban (2005)  | Prospective cohort      | 389          | Pre- and postoperative LMWH       | Enoxaparin 40 mg q12h              | NR                           | 0.6             | 0.26                | NR            |
|                          |                         | 180          | Postoperative LMWH                | Enoxaparin 40 mg q12h              | 0.5–1.5 (range)               | PE 0.6          | 1.7                 | NR            |
|                          |                         | 84           | Postoperative LMWH                | Enoxaparin 40 mg q24h              | 0.5–5 (range)                 | PE 1.2          | 0                   | NR            |
|                          |                         | 180          | Postoperative LMWH                | Enoxaparin 40 mg q24h              | 0.5–1 (range)                 | PE 0            | 1.7                 | NR            |
|                          |                         | 100          | Preoperative LMWH                 | Enoxaparin 30 mg once              | NA                           | PE 2            | 0                   | NR            |
|                          |                         | 124          | Postdischarge LMWH                | Enoxaparin 30 mg q24h              | 10                            | PE 1.6          | 0.8                 | 1.6           |
| Chlysta et al (2014)     | Retrospective cohort    | 192          | Pre- and postoperative LMWH       | Enoxaparin 40 mg (preoperative), then enoxaparin 1 mg per BMI unit (rounded to nearest 10 mg) q12h (postoperative) | Preoperative to discharge | PE 0.5         | Reoperation for bleeding in 1.3%; incidence in each group | NR            |
| Study                        | Design                | Cohort Size | Preoperative Treatment | Postoperative Treatment | Postoperative Duration | Complications |
|-----------------------------|-----------------------|-------------|------------------------|-------------------------|------------------------|---------------|
| Borkgren-Okonek et al (2008) | Prospective cohort    | 124         | Preoperative UFH       | Postoperative and postdischarge LMWH | UFH 5,000              | Preoperative to 10 days postdischarge | 0.8  3.2  0 |
| Singh et al (2012)          | Retrospective cohort  | 11          | Pre- and postoperative LMWH | Enoxaparin 30 mg q12h (BMI < 40) | NR                     | 0  0  NR      |
|                             |                       |             |                        |                         |                        |               |
| Ojo et al (2008)            | Prospective cohort    | 59          | Postoperative and postdischarge LMWH | Enoxaparin 40 mg q12h (BMI > 50) | NR                     | 0  3.5  NR    |
| Raftopoulous et al (2008)   | Prospective cohort    | 132         | Pre- and postoperative LMWH | Enoxaparin 40 mg q12h (BMI 50–59) | NR                     | 0  0  NR      |
|                             |                       |             |                        |                         |                        |               |
|                             |                       | 176         | Pre- and postoperative operative and postdischarge LMWH | Enoxaparin 30 mg preoperative 30 mg, then 30 mg q12h until discharge | Preoperative to 10 days after discharge | 4.5  5.3  0  |
Finks et al identified patient- and procedure-related VTE risk factors through analysis of the Michigan Bariatric Surgery Collaborative (MBSC) database of over 27,000 patients undergoing bariatric surgery in 32 different hospitals. From their analysis, this group developed a preoperative risk assessment model to stratify bariatric surgery patients by VTE risk and then validated this using a bootstrap method. The following characteristics were included in the model: procedure type, patient history of VTE, male sex, BMI, age, and operative time > 3 hours. This scheme was able to stratify patients into low-, medium-, and high-risk groups with <1%, 1–4%, and >4% 30-day VTE event rates, respectively, although approximately 97% of patients were stratified to the low risk group. Data have not yet been published regarding outcomes using this calculator and any corresponding varied approaches.

### Evaluation of strategies to prevent VTE in bariatric surgery patients

In this section, the literature evaluating different methods of VTE prevention in bariatric surgery patients is reviewed, including mechanical and pharmacologic approaches. Studies in which VCF were used as primary prophylaxis against PE are also reviewed.

### Early ambulation

Early postoperative ambulation is reported as a VTE prevention strategy in the included studies, but is not analyzed for effectiveness in isolation of other methods. The purpose of early ambulation is to reduce venous stasis and thereby mitigate the risk of DVT. Early ambulation was included as an adjunctive method of prophylaxis is most of the studies in this review.

### Lower extremity compression

LEC is also believed to reduce the risk of DVT by decreasing venous stasis. LEC may be provided by elastic stockings (ES), such as graduated compression stockings or by intermittent pneumatic compression (IPC) as with sequential compression devices (SCD).

The search criteria used in this review did not identify any studies comparing LEC to no LEC in the bariatric surgery population; most of the included studies applied some form of LEC. However, LEC has been evaluated in other surgical populations, including general surgery. In a Cochrane review, Sachdeva et al reported a 65% reduction in the risk of postoperative DVT in trials in which graduated compression...
### Table 4 Summary of characteristics and outcomes of included studies of vena cava filters for prevention of venous thromboembolism in bariatric surgery patients

| References          | Design                                      | Patients (N) | Intervention | VCF indication(s)                                                                 | Additional VTE prophylaxis                                                                 | VTE events (%) | Bleeding events (%) | Mortality (%) | Filter-related complications (%) |
|---------------------|---------------------------------------------|--------------|--------------|----------------------------------------------------------------------------------|------------------------------------------------------------------------------------------|----------------|---------------------|---------------|----------------------------------|
| Li et al (2012)     | Retrospective cohort                        | 322          | VCF          | NR                                                                                | Variable – 89.8% received anticoagulant                                                  | PE 0.31       | NR                  | 0.31*         | NR                                             |
|                     |                                              | 96,806       | No VCF       | NA                                                                               | Variable – 80.6% received anticoagulant                                                  | PE 0.12       | NR                  | 0.03*         | NA                                             |
| Birkmeyer et al (2013) | Propensity-matched cohort study            | 1,077        | VCF          | NR                                                                                | Preoperative heparin (60% given LMWH) Postoperative heparin (70% given LMWH) Postdischarge heparin (72% given LMWH) | PE 0.84       | NR                  | 0.7*          | 0.6                                            |
|                     |                                              | 1,077        | No VCF       | NR                                                                                | Preoperative heparin (54% given LMWH) Postoperative heparin (68% given LMWH) Postdischarge heparin (66% given LMWH) | PE 0.46       | NR                  | 0.1*          | NA                                             |
| Obeid et al (2007)  | Retrospective cohort                        | 246          | VCF          | Poor mobility, history of VTE, venous disease, BMI > 60, history of VCF            | SCD, “prophylactic” enoxaparin, and warfarin 1 mg/d                                      | PE 0.8        | NR                  | 0.81          | NR                                             |
|                     |                                              | 1,847        | No VCF       | NA                                                                                | SCD and “prophylactic” enoxaparin                                                      | PE 0.59       | NR                  | 0.22          | NA                                             |
| Halmi and Kolesnikov (2007) | Prospective cohort                    | 27           | VCF          | History of VTE, BMI > 65, hypercoagulable state, severe sleep apnea, inability to ambulate, pulmonary hypertension, lower extremity lymphedema | Early ambulation and IPC and preoperative UFH 5,000 U, then 5,000 U q8h or preoperative enoxaparin 40 mg, then 40 mg q12h If high VTE risk also given enoxaparin 40 mg daily for 3 weeks postdischarge | PE 0          | NR                  | 0             | 11.1 (3 minor complications)                  |
|                     |                                              | 625          | No VCF       | NA                                                                                | Early ambulation and IPC and preoperative UFH 5,000 U then 5,000 U q8h or preoperative enoxaparin 40 mg then 40 mg q12h If high VTE risk also given enoxaparin 40 mg daily for 3 weeks postdischarge | PE 0.32       | NR                  | 0             | NA                                             |
| Overby et al (2009) | Prospective cohort                         | 160          | VCF          | Thrombophilia, poor ambulation, history of severe venous stasis, pulmonary hypertension, severe sleep apnea with obesity hypoventilation, history of VTE, BMI > 60 | SCD and UFH 5,000–7,500 U q8h until discharge                                           | PE 0.63       | NR                  | 0.9           | 2.5                                            |

*Continued*
stockings were evaluated as VTE prophylaxis. In a meta-analysis of studies comparing IPC with no prophylaxis, the application of IPC appeared to reduce the risk of DVT by approximately 60%. Complications of ES may include skin breaks, ulcers, blisters, and skin necrosis.

The favorable risk profile and potential benefits of LEC support its use in postoperative bariatric surgery care.

### Pharmacologic prophylaxis

In bariatric surgical practice, pharmacologic prophylaxis is commonly used. In this section, evaluations of specific medications, their dosing, and duration of use are presented. In these evaluations, the effectiveness of prophylaxis and its safety are both considered since anticoagulants are protective against VTE but have the potential to increase postoperative bleeding.

### Comparison of either UFH or LMWH with no pharmacologic prophylaxis

Direct comparisons of heparins with no pharmacologic VTE prophylaxis after bariatric surgery are limited. Thus, it is useful to consider data from general surgery populations. Multiple studies have evaluated UFH and LMWH for VTE prevention in general surgery and have been the subject of previous reviews. Trials of UFH in surgical patients, including general surgical patients, show risk reductions of 47% and 41% for fatal and nonfatal PE, respectively. Data from eight clinical trials in general surgery, in which LMWH was compared with placebo or no prophylaxis, suggest a 71% risk reduction of overall clinical VTE events and PE (RR, 0.29; CI, 0.11–0.73). These reviews also show that...
Multiple studies have compared UFH with LMWH for VTE prevention in general surgery patients. In a meta-analysis by Mismetti et al, there was no statistically significant difference between UFH and low-dose LMWH (3,400 anti-Xa units) with respect to clinical PE or DVT outcomes; high-dose LMWH (>3,400 anti-Xa units) was associated with a lower risk of clinical PE, but an increased risk of major hemorrhage.

Only two included studies compared UFH with LMWH for VTE prevention in bariatric surgery. Using the MBSC database, Birkmeyer et al compared three prophylaxis regimens; preoperative UFH and postoperative UFH, preoperative UFH and postoperative LMWH, and both pre- and postoperative LMWH, with the former group serving as reference. SCD were used by 98% of patients and 3.2% also received a VCF. The groups were compared with mixed-effects logistic regression. Overall, the risk of VTE was 66% lower in groups receiving LMWH compared with the group receiving postoperative UFH, (OR, 0.34; CI, 0.19–0.62). This difference persisted among subgroups which included only patients at low risk of VTE events (defined as <1%). However, when only patients at high risk of VTE events (≥1%) were analyzed, the risk of VTE was not statistically significantly different in groups receiving LMWH compared to UFH (OR, 0.37; CI, 0.19–1.22). Of note, medication doses and duration are not described in this analysis. There was no significant difference across groups with respect to serious hemorrhage, defined as transfusion of >4 units of blood product or reoperation for bleeding. Mortality was not reported in this study.

In a nonrandomized study of two consecutive cohorts of 238 patients each, Kothari et al compared enoxaparin 40 mg SC twice daily to UFH 5,000 units SC three times daily. All patients received SCD and early ambulation. All underwent laparoscopic RYGB. There were no DVT cases in either group, and only one PE was seen in the UFH group. The average operative time was longer in the UFH group compared to the LMWH group (160 vs 129.5 minutes), but the average BMI was slightly higher in the LMWH group. Postoperative transfusion was given more frequently among the LMWH group compared to the UFH group (5.9% vs 1.3%, P=0.011), and four patients in the LMWH group required reoperation for bleeding. The dose of enoxaparin was higher than standard, potentially influencing the bleeding observations.

The data comparing UFH with LMWH prophylaxis in bariatric surgery are also limited and do not indicate...
superiority of one over another with respect to VTE and bleeding complications. Data from the two described studies do not allow for comparisons of standard heparin doses. Until such data are available, use of either UFH or LMWH appears satisfactory.

Evaluation of adjusted-dose heparin

Optimal dosing of prophylactic heparin in obese patients such as those undergoing bariatric surgery is unclear. For example, therapeutic LMWH dosing is calculated from total body weight, raising the question as to whether standard prophylactic doses of LMWH are sufficient protection against VTE in obese patients.

Indeed, the majority of studies of pharmacologic prophylaxis in this review evaluated adjusted-dose LMWH after bariatric surgery. In a multicenter pilot study of two doses of LMWH for prevention of VTE in bariatric surgery, Imberti et al randomized 250 subjects to receive either nadroparin 4,250 IU/d (standard prophylactic dose) or 6,400 IU/d (150% of standard dose). Block randomization was stratified by center, patient sex, and BMI. Study medication was given preoperatively and for 9±2 days postoperatively, and ascertainment of VTE was conducted only during the time subjects received medication. The groups did not differ with respect to sex, age, BMI, operative time, or other VTE risk factors. The rates of VTE among the standard and adjusted-dose groups were 1.5% (CI, 0.2–6.0) and 0.8% (CI, 0.4–5.3), respectively, and were not statistically significantly different. The rates of combined major bleeding and clinically relevant nonmajor bleeding were 6.1% (CI, 2.9–12.1) and 5% (CI, 2.1–11.1) for the standard and adjusted-dose groups, respectively. Major bleeding was not reported separately. Being a pilot study, this analysis may not have been adequately powered to detect true difference in effect sizes. In the only other randomized study of pharmacologic prophylaxis in bariatric surgery, Kalfarentzos et al compared nadroparin 5,700 IU SC daily (standard prophylactic dose) to 9,500 IU SC daily (167% of standard dose) in 60 patients undergoing RYGB. There were no VTE events in either group. Given the current knowledge of typical VTE event rates, the small sample size likely did not allow a valid comparison of efficacy. There were two subjects in the higher dose group who developed major bleeding compared with none in the standard-dose group.

Two studies evaluated increased LMWH dosing not based on patient BMI. Scholten et al analyzed results from two consecutive groups of patients who were administered different enoxaparin dosing schedules, 30 mg SC twice daily (92 patients), and 40 mg SC twice daily (389 patients). Most patients underwent open RYGB. All patients reportedly received early ambulation and LEC. Patients with previous VTE were offered postdischarge anticoagulation, but the frequency of this is not reported. Patients in the higher dose group had shorter operative times and length of hospital stay; other reported VTE risk factors were not different between groups. The higher dose group had a lower incidence of VTE events (0.6% vs 5.4%, P<0.01), with no significant difference in bleeding. Of note, there were four PEs in the lower dose group and none in the higher dose group. Hamad and Choban compared postoperative VTE rates in five hospitals using different LMWH prophylaxis regimens, including one center at which a single preoperative 30 mg dose of enoxaparin was used and another at which enoxaparin 30 mg once daily was given only beginning at discharge. Other centers used 40 mg enoxaparin doses with either once or twice daily dosing frequency. The centers were heterogeneous with respect to the prevalences of VTE risk factors among their patients, and operative times reportedly varied, making valid comparative analysis difficult.

Several cohort studies report results of adjusted-dose LMWH based on patient BMI. Chysta et al compared three regimens of adjusted-dose enoxaparin with respect to thromboembolism, bleeding, and mortality (Table 3). There were significant differences in age, procedure type, and anastomotic leaks between the groups (the latter being more common in the group receiving enoxaparin 40 mg preoperatively). There were no significant differences in VTE or bleeding rates among groups. One PE occurred in the group receiving fixed preoperative enoxaparin dosing. This individual’s course was complicated by conversion to open surgery, ventral hernia repair, and wound infection requiring debridement. The authors report that 29 patients were excluded from the analysis as they were not managed by the described protocols. It is possible that these patients were managed differently due to differences in perceived VTE or bleeding risk, introducing bias into the results.

Borkgren-Okonek et al compared two different postoperative enoxaparin doses following primary RYGB (93% laparoscopic) depending on patient BMI (BMI ≤50 given enoxaparin 40 mg twice daily during hospitalization, BMI >50 given enoxaparin 60 mg twice daily during hospitalization). All patients received UFH 5,000 units 2 hours preoperatively, IPC, and early ambulation; LMWH was continued once daily for 10 days after discharge. Patients with a known hypercoagulable disorder or a history of VTE were excluded. Antifactor-Xa assays were performed during
LMWH prophylaxis, with 10 mg dose adjustments made for what were deemed levels out of prophylactic range; of 223 total patients, 37 had dose increases administered. Overall, only one patient (in the enoxaparin 40 mg group) had a VTE event; major bleeding events were noted in five patients (four requiring transfusions and one reoperation) with four bleeds in the 40 mg group. Singh et al\textsuperscript{57} also analyzed BMI-adjusted LMWH dosing in a cohort of 170 patients undergoing RYGB. All patients received LMWH, 1 hour preoperatively, IPC, and early ambulation. Postoperatively, enoxaparin was given SC twice daily with the following dosing: BMI $\leq$ 40, 30 mg; for BMI 41–49, 40 mg; for BMI 50–59 and for BMI $> 59$, 60 mg. Overall, there were no VTE events, and five patients had bleeding events, with four of these in the 40 mg group. Of note, 145 of the 170 patients were in the 40 mg group, limiting intergroup comparisons.

Ojo et al\textsuperscript{48} analyzed results from patients who underwent open RYG and received either enoxaparin 40 mg SC twice daily or 60 mg SC twice daily, starting in the postoperative period and continuing for 14 days postdischarge. The administered dose was selected by the attending surgeon. Only patients deemed to be at higher risk of VTE were included (personal history of VTE, BMI $\geq$ 60, or BMI $\geq$ 50 with venous stasis, obstructive sleep apnea, or severe limitation to ambulation). Patients with a history of bleeding or on chronic anticoagulation were excluded. The primary study aim was to ascertain the incidence of major bleeding complications, defined as bleeding during LMWH administration resulting in drug discontinuation, bleeding-related readmission, blood transfusion, or intervention for bleeding. There were no major bleeding events during the 2 week postdischarge study period.

A meta-analysis by Ikesaka et al\textsuperscript{55} evaluated the efficacy and safety of adjusted-dose heparin in patients undergoing bariatric surgery. Several studies discussed earlier were not included in this meta-analysis likely due to publication date, lack of VTE event reporting, or unclear treatment frequencies.\textsuperscript{23,25,27,28,31} These investigators chose to include two additional studies discussed in this review and one that is not.\textsuperscript{20,21,56} The study by Shepherd et al\textsuperscript{56} is a single-arm study of LMWH, adjusted daily to a target antifactor-Xa level of 0.1–0.25 U/mL. In the meta-analysis, the rates of in-hospital VTE were 0.54% (CI, 0.2–1.0) and 2.0% (CI, 0.1–6.4) for 1,428 adjusted-dose and 430 nonadjusted-dose patients, respectively. For major bleeding, the effect sizes were 1.6% (CI, 0.6–3.0) and 2.3% (CI, 1.1–3.9) for those respective groups. Of note, the study by Shepherd et al\textsuperscript{56} contributed approximately half of the patients in the adjusted-dose group. Ikesaka et al\textsuperscript{55} used $F$ as their reported measure of heterogeneity. For the adjusted-dose patient VTE rate, $F$ = 0% and for major bleeding, $F$ = 63.3%, while for non-adjusted-dose patient VTE rate, $F$ = 71.8% and for major bleeding, $F$ = 0%. Variance in effect sizes across studies may be due to either sampling error or to some degree of variance in true effect, depending on the measure and patient group. All $F$ estimates had wide CIs.

Adjustment of LMWH dose using an antifactor-Xa level was used in two aforementioned studies.\textsuperscript{26,56} A detailed discussion regarding the use of antifactor-Xa levels to monitor prophylactic LMWH is beyond the scope of this review, but has been the subject of review elsewhere.\textsuperscript{57,58} There is currently no definitive supportive evidence correlating antifactor-Xa level and postoperative VTE and bleeding risk for bariatric surgery patients.

Concern that standard prophylactic heparin dosing is not optimal in bariatric surgery is reasonable and deserves continued investigation. Currently available data do not support specific strategies for adjusted-dose heparin for VTE prevention in this group. A common design limitation is the lack of comparison of different strategies within groups of patients of similar risk or BMI.

Evaluation of postdischarge heparin prophylaxis

VTE events may occur in the immediate postoperative period or after hospital discharge. Based on clinical trials, some guidelines endorse consideration of postdischarge prophylaxis in abdominal or pelvic cancer surgery and in major orthopedic surgery.\textsuperscript{51,59} Postdischarge VTE after bariatric surgery is common.\textsuperscript{57} For example, Froehling et al\textsuperscript{8} found that the incidence of VTE rose from 0.3% to 1.9% between 7 and 30 days postoperatively. These observations raise the question as to whether the duration of VTE prophylaxis should be extended for bariatric surgery patients.

Two of the included studies specifically evaluated postdischarge pharmacologic prophylaxis in the bariatric surgery population.\textsuperscript{20,21} Other studies in this review included extended duration anticoagulation, but did not compare this practice to standard duration prophylaxis.\textsuperscript{33,26,28} Raftopoulos et al\textsuperscript{50} compared administration of enoxaparin 30 mg twice daily until hospital discharge only (132 patients) and the addition of enoxaparin 40 mg daily for 10 days following discharge (176 patients). The former group also received enoxaparin 30 mg given 1 hour preoperatively. Of the group who received in-hospital prophylaxis only, 4.5% experienced a VTE event within 30 days of surgery (three patients with PE and three
with DVT) compared with none in the post-discharge group ($P=0.006$ for the comparison). Four of the six events in the hospital-only prophylaxis group occurred between hospital discharge and 30 days postoperatively. There was no significant difference in bleeding events between the groups (0% vs 4.5%, $P=0.06$). One patient in each group required reoperation for bleeding. There were no deaths in either group.

Cossu et al$^{13}$ compared patients receiving UFH once prior to anesthesia with patients receiving UFH 4–5 days preoperatively and 8–9 days postoperatively, with doses based on activated partial thromboplastin time monitoring, followed by low-dose UFH for at least 15 days after discharge. The authors found a trend toward fewer PE in the extended duration protocol group (1.2% vs 3%, $P =$ not significant) but higher rates of bleeding (2.3% vs 0% requiring transfusions, $P =$ not significant). This study essentially compares postoperative UFH prophylaxis with no pharmacologic prophylaxis, making the impact of postdischarge anticoagulation uncertain.

Analysis of postdischarge pharmacologic prophylaxis for bariatric surgery has been limited, but results from Raftopoulos et al$^{10}$ are promising. Given what is known regarding the timing of postbariatric surgery, longer duration prophylaxis of VTE events merits further evaluation.

**Evaluation of oral anticoagulants as VTE prophylaxis**

Oral anticoagulants such as warfarin and other vitamin K antagonists, direct thrombin inhibitors (dabigatran), and factor Xa inhibitors (rivaroxaban, apixaban, and edoxaban) have been evaluated for VTE prevention after orthopedic surgery, but not in general surgery.$^{60,61}$ We did not identify any studies of oral direct thrombin inhibitors or factor Xa inhibitors for prevention of VTE in bariatric surgery patients. Heffline$^{12}$ described a comparison of a VTE rates in an initial cohort given a combination of UFH and SCD, with a subsequent cohort given a similar regimen except the addition of postoperative warfarin (INR [international normalized ratio] goal of <1.8). Whether the two cohorts differed with respect to the types of procedures or patient characteristics was not reported. After implementation of the warfarin protocol, VTE events appeared to decrease; statistical significance was not reported. Of note, in the warfarin recipient cohort, patients deemed high risk received VCF, but the number of patients with VCF is not reported. Bleeding events were also not reported.

Given the limited data, oral anticoagulants, including warfarin, are not recommended for VTE prophylaxis in bariatric surgery patients. Neither their efficacy nor associated bleeding risk has been adequately investigated in this patient population.

**Evaluation of VCFs as VTE prophylaxis**

VCF for primary prevention of PE in bariatric surgery patients has been investigated in six studies identified in this review.$^{15,16,22,29,30,33}$

Li et al$^{15}$ reported the outcomes of 322 patients in the Bariatric Outcomes Longitudinal Database (BOLD) who had preoperative placement of a VCF with those ~97,000 patients who did not. The presence of multiple patient and procedural characteristics favoring higher risk of VTE were seen with greater frequency in the VCF group, and they were more likely to receive anticoagulation and SCD. Results showed higher rates of DVT (0.93% vs 0.12%, $P=0.001$) but no statistically significant difference in PE (0.31% vs 0.12%) when comparing VCF recipients with those who did not have a VCF respectively. All-cause mortality was also higher in the VCF group (0.31% vs 0.03%, $P=0.003$). Although improved outcomes were not associated with VCF use, the study design allows the possibility that VCF recipients were high-risk patients whose postoperative PE rate would have been higher without the addition of a VCF.

Birkmeyer et al$^{16}$ found similar results in an analysis from the MBSC database. Using data from patients who underwent bariatric surgery between 2006 and 2012, Birkmeyer et al$^{16}$ identified 1,077 propensity matched controls for 1,077 patients who received VCF. These groups did not differ with respect to patient or procedure characteristics, although the VCF group was more likely to receive preoperative LMWH and postdischarge LMWH. Results showed a significantly higher risk of DVT (1.2% vs 0.4%; OR, 3.3; $P=0.039$) in the VCF group and a nonsignificant trend toward higher mortality (0.7% vs 0.1%; OR, 7.0; $P=0.068$). There was also a nonsignificant trend toward increased risk of PE in the VCF group (0.84% vs 0.46%; OR, 2.0; $P=0.232$).

In another registry study, Obeid et al$^{33}$ reported nonsignificant trends toward higher rates of PE (0.8% vs 0.59%), DVT (1.21% vs 0.65%), and mortality (0.81% vs 0.22%) in 246 patients who received a VCF compared to those who did not. VCF recipients were more likely male and had a higher average BMI, but the distribution of procedure type did not differ between groups. Indications for VCF included previous VTE, poor mobility, venous disease, and BMI $>60$ kg/m$^2$. Despite the selection of higher risk patients for VCF, outcomes did not appear to improve with this intervention.
although again, it is unknown whether PE rates would have been higher without VCF.

In contrast, three studies reported more favorable outcomes with VCF. Halmi and Kolessnikov29 found a nonsignificant trend toward lower rates of DVT and PE in patients receiving VCF (0% vs 0.32% and 0% vs 7.12%, respectively). Their study reports outcomes for 652 patients undergoing mini-open RYGB. Of these patients, 27 received a VCF based on indications of previous VTE, BMI >65, severe sleep apnea, a hypercoagulable state, pulmonary hypertension, lower extremity lymphedema, or inability to ambulate. However, the study reported VCF patients deemed to be at “significant VTE risk” (indications not reported) were given an additional 3 weeks of antithrombotic therapy. Overby et al30 published an experience with 330 patients undergoing primary RYGB in which high risk patients received a VCF. High risk was defined as the presence of a thrombophilia, BMI >60, history of VTE, severe venous stasis or severe sleep apnea with obesity-hypoventilation, poor ambulation, or pulmonary hypertension. In this analysis, VCF recipients had lower rates of PE (0.63% vs 2.94%) but higher rates of DVT (3.13% vs 2.35%). Neither of these differences reached statistical significance.

Gargiulo et al22 describe results of three practice patterns for VTE prevention after open RYGB (Table 4). In a retrospective review, they noted that PE occurred in patients without VCF who had BMI >55. In the subsequent period in which BMI >55 was added to the indications for VCF placement, no patients with BMI >55 and a VCF had postoperative PE. In a third period during which patients with a BMI of >55 were offered optional VCF placement, the incidence of PE (0% vs 28%, P<0.05) and PE-specific mortality (0% vs 11%, P<0.05) appeared to be less with VCF. These small groups did not appear to differ on other reported risk factors for VTE, but the process by which patients opted in for VCF placement was not described.

Complications directly related to VCF include strut fracture, filter migration, vena cava perforation, VCF thrombosis, and insertion site infection or thrombosis.52 VCF migration may involve serious injury to heart valves, the myocardium, or pericardium. Although uncommon, understanding the risk of these complications is important when VCF placement is being considered. Among the included studies, four reported VCF-related complications.16,22,29,30 For example, in the analysis by Overby et al,30 complications included pneumothorax, filter migration to the right atrium, hemopericardium, and inability to ablate an accessory pathway in atrial fibrillation due to the filter. In their analysis, Birkmeyer et al16 reported VCF complications including a damaged heart valve requiring replacement, two fatal inferior vena cava thrombi, contrast nephropathy, and an incision site infection. Retrieval of VCF was reported in two studies, with success rates >90% in both.29,30

There have been two published systematic reviews with meta-analyses addressing the use of VCF in bariatric surgery.63,64 Both included the studies discussed earlier with the exception of the study by Hamli and Kolessnikov,29 which was not included by Brotman et al.63 Both meta-analyses indicate that VCF do not appear to decrease the risk of PE. The RR of PE determined by Kaw et al64 was 1.02 (CI, 0.31–3.37) and by Brotman et al63 1.21 (CI, 0.57–2.56). VCF did appear to increase the risk of postoperative DVT by both meta-analyses; Brotman et al63 found a RR of 2.94 (CI, 1.35–6.38) and Kaw et al64 a RR of 2.81 (CI, 1.33–5.97). Brotman et al63 found a significantly increased risk of mortality among recipients of VCF across their included studies (RR, 4.30; CI, 1.60–11.54), while Kaw et al64 noted a nonsignificant trend toward higher mortality rates among VCF recipients (RR, 3.27; CI, 0.78–13.64). Studies by Hamli and Kolessnikov29 and Overby et al30 were not included in the former mortality analysis.

In the meta-analysis by Kaw et al64 the only reported measure of heterogeneity is I². For the estimated effect sizes, I² ranged from 35% to 60%, suggesting that a proportion of the observed variance in effect sizes may reflect true differences across studies. Brotman et al63 found a similar I² result for DVT risk ratio (40.3%), but for PE and mortality effect sizes the I² were 6.9% and 0%, respectively, suggesting that for those measures, most or all of the observed variance is due to sampling error.

Available data do not appear to support the routine placement of VCF as an adjunctive method of prophylaxis in bariatric surgery. Currently, all the available data on VCF placement is observational in nature, which limits the ability to accurately determine efficacy of an intervention; randomized controlled trials are needed for this.

Selected published guideline recommendations

Several organizations or professional societies have published guideline recommendations for prevention of VTE in bariatric surgery patients (Table 6).51,65–67 None of the published guidelines reviewed discusses different approaches to VTE prevention based on procedure type, BMI, or open versus laparoscopic surgeries. Guidelines do reflect the uncertainties in the literature discussed.
| Guidelines (year)                                                                 | Early ambulation | Lower extremity compression | Pharmacologic prophylaxis                              | Adjusted-dose heparin | Postdischarge pharmacologic prophylaxis | Prophylactic vena cava filter | Other recommendations                                                                 |
|--------------------------------------------------------------------------------|------------------|----------------------------|------------------------------------------------------|-----------------------|----------------------------------------|-------------------------------|--------------------------------------------------------------------------------------|
| American Association of Clinical Endocrinologists/ The Obesity Society/ American Society for Metabolic and Bariatric Surgery (2013) | Recommended      | Recommended SCD            | SC UFH or LMWH given within 24 hours of surgery.     | Not mentioned         | Consider for high risk patients (history of DVT) | Risk may exceed benefit        | Discontinue estrogen therapy preoperatively. Patients with history of DVT or cor pulmonale should undergo evaluation for DVT |
| American Society for Metabolic and Bariatric Surgery (2013)                     | Recommended      | Recommended for all        | Combination of mechanical and chemical, should be considered based on clinical judgment and risk of bleeding. Conflicting data but data suggest LMWH over UFH | Not mentioned         | Consider but insufficient data to recommend specific dose or duration | VCF as only method not recommended. Consider addition of VCF in high risk patients where VTE risk > risk of filter-related complications | None                                                                                |
| American College of Chest Physicians (2012)                                     | Recommended      | Rogers or Caprini score recommended: Low VTE risk: IPC Moderate VTE risk: Not high bleeding risk: SC UFH or LMH or IPC High bleeding risk: IPC High VTE risk: Not high bleeding risk: SC UFH or LMWH and ES/IPC High bleeding risk: IPC | Not mentioned         | Not mentioned         | Not recommended                       | If heparins contraindicated and not high risk of bleeding, consider low dose aspirin, fondaparinux or IPC | None                                                                                |
| Interdisciplinary European Guidelines on Metabolic and Bariatric Surgery (2013)  | Recommended      | Recommended ES and IPC     | SC LMWH                                             | Not mentioned         | Not mentioned                         | Not mentioned                 | None                                                                                |

**Abbreviations:** SCD, sequential compression devices; SC, subcutaneous; UFH, unfractionated heparin; LMWH, low-molecular-weight heparin; DVT, deep vein thrombosis; VCF, vena cava filters; VTE, venous thromboembolism; IPC, intermittent pneumatic compression; ES, elastic stockings.
The American Association of Clinical Endocrinologists, the Obesity Society, and the American Society for Metabolic and Bariatric Surgery (AACE/TOS/ASMBS) together have produced guidelines for bariatric surgery.69 Early ambulation and IPC as well as postoperative UFH or LMWH are recommended. Anticoagulant dosing is not specified, but postdischarge pharmacologic prophylaxis is recommended for “high-risk” patients, such as those with history of DVT; no specific duration of therapy is suggested. These organizations discuss that VCF may present a greater risk than benefit due to filter-related complications. They also recommend preoperative discontinuation of estrogen medications (one cycle of oral contraceptives and 3 weeks for hormone replacement therapy) since these may increase the risk of VTE. Finally, they recommend preoperative DVT screening for patients with a history of DVT or cor pulmonale; evidence supporting this consideration is not discussed.

The ASMBS provides a separate set of recommendations.68 The ASMBS recommends prophylaxis with a combination of early ambulation and mechanical prophylaxis for all patients. They state that use of pharmacologic prophylaxis “should be considered based on clinical judgment and risk of bleeding”. The ASMBS expresses a preference for LMWH over UFH, although they stipulate that there is conflicting data regarding the type of pharmacologic prophylaxis to use. The ASMBS also recommends extended duration of pharmacologic prophylaxis but do not provide specific dose or duration recommendations. Although VCF are not recommended as the only method of prophylaxis, addition of VCF to mechanical and pharmacologic prophylaxis may be considered in selected high-risk patients for whom the risk of VTE outweighs the risk of filter-related complications.

The American College of Chest Physicians (ACCP) does not offer specific recommendations for bariatric surgery patients, but includes this group with patients having other abdominal, vascular, or plastic reconstructive surgery.51 The ACCP recommends using either a Rogers et al35 or Caprini34 score to stratify patients as low, moderate, or high risk of VTE. Given usual body habitus and type of surgery, bariatric surgery patients typically will be considered moderate to high VTE risk, depending on comorbidities. The ACCP also recommends stratifying bleeding risk to determine a prophylaxis plan. For patients with a moderate VTE risk who are not considered to be at high bleeding risk, LMWH, UFH, or IPC may be used; for moderate VTE risk but high bleeding risk, IPC alone is recommended. For patients with a high VTE risk, a combination of mechanical (IPC or ES) and pharmacologic (either LMWH or UFH) measures are recommended, unless the bleeding risk is high (IPC alone is recommended). The ACCP also recommends against the use of VCF for primary prophylaxis and does not offer specific recommendations regarding adjusted dose heparins or extended duration anticoagulation postdischarge.

The Interdisciplinary European Guidelines on Metabolic Surgery recommend VTE prevention for all bariatric patients through LMWH administration, use of LEC (both ES and SCD), and early postoperative ambulation.67 These guidelines do not address the questions of augmented LMWH dosing, postdischarge anticoagulation, or VCF placement.

Published guidelines have in common the recommendation for early ambulation and LEC and generally concur regarding the use of heparin prophylaxis. However, the uncertainties regarding adjusted-dose and post-discharge heparin are reflected in the guidelines. They are also generally concurrent in their recommendations regarding prophylactic VCF.

Conclusion and summary recommendations

VTE continues to be an important source of postoperative morbidity and mortality among patients undergoing bariatric surgery, despite current VTE prevention methods. The practice of postbariatric surgery VTE prophylaxis has been primarily supported by data from the general surgery literature. Postoperative care encouraging early ambulation, use of LEC, and pharmacologic prophylaxis (assuming satisfactory bleeding risk) appears prudent. To date, the literature regarding further optimization of preventive approaches specific to bariatric surgery patients has been limited. However, the principles behind adjusted-dose heparin and post-discharge prophylaxis are worthy of further analysis with randomized controlled trials to assess their efficacy and safety. At present, VCF do not have an established role in bariatric surgery VTE prophylaxis. Studies are also needed which better control for patient and procedure-related VTE risk factors, and future studies may incorporate what is known regarding patient and procedure-related risk factors to develop validated, stratified management plans using different intensities of prophylaxis. Until more data are available, institutional quality improvement efforts should focus on ensuring consistent application of established methods.

Disclosure

The authors report no conflicts of interest in this work.
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