ABSTRACT
The shortcomings of current anticoagulants have led to the development of newer, albeit more expensive, oral alternatives.

Objective: To explore the potential impact the new anticoagulants dabigatran and rivaroxaban in the local hospital setting, in terms of utilisation and subsequent costing.

Method: A preliminary costing analysis was performed based on a prospective 2-week clinical audit (29th June – 13th July 2009). Data regarding current anticoagulation management were extracted from the medical files of patients admitted to Ryde Hospital. To model potential costing implications of using the newer agents, the reported incidence of VTE/stroke and bleeding events were obtained from key clinical trials.

Results: Data were collected for 67 patients treated with either warfarin (n=46) or enoxaparin (n=21) for prophylaxis of VTE/stroke. At least two-thirds of all patients were deemed suitable candidates for the use of newer oral anticoagulants (by current therapy: warfarin: 65.2% (AF), 34.8% (VTE); enoxaparin: 100%, (VTE)). The use of dabigatran in VTE/stroke prevention was found to be more cost-effective than warfarin and enoxaparin due to significantly lower costs of therapeutic monitoring and reduced administration costs. Rivaroxaban was more cost-effective than warfarin and enoxaparin for VTE/stroke prevention when supplier-rebates (33%) were factored into costing.

Conclusion: This study highlights the potential cost-effectiveness of newer anticoagulants, dabigatran and rivaroxaban, compared to warfarin and enoxaparin. These agents may offer economic advantages, as well as clinical benefits, in the hospital-based management of anticoagulated patients.

Keywords: Anticoagulants. Costs and Cost Analysis. Australia
INTRODUCTION

Arterial and venous thromboses remain the major cause of mortality and morbidity in the developed world. While arterial thrombosis has been directly associated with the development of myocardial infarction and stroke, venous emboli originate from deep vein thrombosis in more than 95% of cases.2 These emboli are transported across large channels and generally pass through the right side of the heart and into the pulmonary vasculature to form fatal, pulmonary emboli.3 The coagulation pathway plays a pivotal role in the formation and progression of both arterial and venous thrombi. Therefore, drugs aimed at inhibiting this pathway are a major stratagem for the development of novel antithrombotic agents.6-8

Traditional anticoagulants such as warfarin, unfractionated heparin (UFH), and low-molecular weight heparins (LMWH) (e.g., enoxaparin, dalteparin, tinzaparin) have dominated the prevention and treatment of thromboembolic conditions such as myocardial infarction, stroke, and post-surgical thrombosis.5,8 While heparins are quick-acting intravenous or intramuscular preparations used over short periods in the hospital setting, warfarin is an oral Vitamin K antagonist that is suitable for long-term therapy.10-13 Although it is the only oral anticoagulant currently approved for widespread use, warfarin carries a number of limitations including: (1) slow onset of action; (2) the potential for genetic variation in metabolism; (3) multiple food and drug interactions; and (4) a narrow therapeutic index. Consequently, new “improved” anticoagulants, which demonstrate similar efficacy as warfarin but importantly with a better safety profile with respect to bleeding are highly sought-after.6-8

Selective newer, oral, anticoagulants which act on a single coagulation factor in the coagulation cascade have been recently developed. These include direct thrombin inhibitors such as dabigatran (Pradaxa®), and the factor Xa inhibitor, rivaroxaban (Xarelto®). In contrast to the conventional multi-targeted antithrombotics such as warfarin and heparins, these agents have the potential to be much more effective, safer, and easier to use.6-9

Dabigatran is an oral thrombin inhibitor that inactivates thrombin directly by binding to thrombin’s active site and preventing its interaction with substrates. The drug offers a rapid onset of action compared to traditional anticoagulants.14,15 A phase II ‘efficacy and safety’ study has highlighted it’s therapeutic benefits in prophylaxis of venous thromboembolism in patients undergoing orthopaedic surgery, where gastrointestinal complaints represented the most frequent adverse event.5,8 Dabigatran has also shown superiority over enoxaparin in two large phase III trials for prophylaxis of venous thromboembolism following total knee replacement and in patients undergoing total hip replacement.16,15 Moreover, dabigatran demonstrates that fixed doses of oral anticoagulants can indeed be used without the necessity for routine coagulation monitoring.6

Rivaroxaban is another selective inhibitor of factor Xa that has currently been approved for the prevention of thromboembolism in adult patients undergoing elective hip or knee replacement surgery.6-9,14-19 Traditionally, LMWHs (e.g., enoxaparin) and warfarin have been prescribed for the prevention of thromboembolism following orthopaedic surgery.5-9 However, intravenous/subcutaneous administration of LMWHs and the reduced safety profile for warfarin limit the ongoing use of these agents, making this alternative oral therapeutic a more convenient option for both the patients and clinicians. Rivaroxaban represents a highly potent and competitive factor Xa inhibitor that can inhibit fibrin-bound factor Xa with a significantly greater efficacy than fondaparinux when used at similar concentrations.20 In three phase III multicentre, double-blinded double dummy, randomised controlled trials, oral rivaroxaban 10mg demonstrated similar efficacy with intravenous enoxaparin 40mg.21 Additionally, the incidence of side effects did not differ between treatment groups in the same study.21

Direct thrombin inhibition, and inhibition of factor Xa represent novel approaches for the prevention and treatment of thromboembolism.6-9,16-19 However, their role in the management of hospitalised patients is unclear in terms of potential indications and the impact of this on hospital treatment costs, given that these agents are more expensive than those currently used.16-19 Therefore, the aim of this study was to evaluate the potential impact of the use of the new anticoagulants dabigatran and rivaroxaban in the local hospital setting for the prevention of venous thromboembolism (VTE) and stroke in patients with atrial fibrillation (SPAF), and subsequent estimate the costs involved.

METHODS

Data Collection

To evaluate the potential impact of dabigatran and rivaroxaban use in the local hospital setting, the costing analysis was designed to consider acute and long-term VTE and SPAF events, their long-term effects, prophylaxis-related side effects (eg. bleeding, deaths associated with VTE and SPAF), and other adverse reactions. A descriptive study was undertaken at a teaching hospital (Ryde Community Hospital, 170-beds) within the Sydney Northern Area Health Service comprising a prospective clinical audit conducted over a two week period (29th June -13th July 2009). The data collected was then used to undertaken a preliminary pharma-economic evaluation of the potential impact of using these agents in the hospital setting. A simplified cost-effectiveness analysis was undertaken to compare the costs and clinical outcomes of rivaroxaban, dabigatran, warfarin and enoxaparin for the prevention of VTE and stroke. Approval was obtained from the institutional Human Research Ethics Committee.
Using purpose-designed forms, prospective data was collected on patients aged 18 years and above who had been admitted under the Geriatric Medicine, Orthopaedics and Cardiology wards and had been receiving either warfarin or enoxaparin during their hospital admission. Medical notes and drug charts were screened manually to identify patients on warfarin or enoxaparin and then to identify patients where oral rivaroxaban or dabigatran may be indicated. Data were also collected on patient characteristics, e.g. age, and gender, indications for anticoagulation, details of therapy, blood clotting test results (e.g. International Normalised Ratio, INR), incidence and severity of bleeding events, risk factors for adverse events including drug interactions and comorbidities.

**Decision Model**

The principal outcome measure was the incremental cost gained free from total venous thromboembolism (VTE) and stroke in patients with atrial fibrillation (SPAF). This end point was applied as the primary efficacy model in this study and was used to estimate the probability of VTE and stroke in patients, following orthopaedic surgery (hip and knee), and with atrial fibrillation, respectively (Figure 1A). A VTE could include proximal or distal DVT and could be symptomatic or non-symptomatic. A stroke could be symptomatic or immediately fatal. Symptomatic events were either detected by the patient at the hospital or were detected following discharge and therefore required readmission and/or outpatient treatment. Patients could also present with symptoms of VTE/stroke that were not confirmed, therefore resulting in additional cost increment associated with diagnostic procedures but had no significant effect on the patient outcome. Patients could also die following surgery from causes that were unrelated to either VTE or stroke (other post-surgical death).

Major bleeding events were incorporated into the study and included gastrointestinal, intracranial, or bleeding at surgical site so that the costs associated with these events could be accounted for. Patients with major bleeding could recover, die or be permanently disabled. Patients with minor bleeding events were assumed to recover (Figure 1B).

![Figure 1: Schematic diagram of the model structure: (A) Model for VTE/SPAF; (B) Diagram of adverse reactions](image-url)
Event Probabilities

Data from subjects were used for the number of VTE and SPAF due to warfarin and the number, type, and severity of bleeding events. The treatment of major bleeding events was modelled from review of the literature and data from the prospective audit. For dabigatran, the probability of VTE and/or SPAF, major and/or minor bleeding events were determined based on incidence of events in the total knee replacement (TKR) trial (RE-MODEL) and the total hip replacement (THR) trial (RE-NOVATE). For rivaroxaban, the probabilities of total VTE and/or SPAF and all-cause mortality, major and minor bleeding were calculated from the RECORD/PETRO trial. Probabilities for the new oral anticoagulants were quantified by assessing the relative risk for each end point, compared to the probabilities in the warfarin and enoxaparin arm.

Patient Population

The impact of new anticoagulants were modelled separately to reflect differences in the incidence of venous thromboembolism (VTE), and stroke in patients with atrial fibrillation (SPAF), age and gender distribution, and duration of VTE and SPAF prophylaxis. The mean age of patients incorporated into the model was 81.5 years for VTE and 83.5 years for SPAF. Subgroup analysis was performed by age, gender, and risk factors for VTE and SPAF.

Resource Use and Cost Estimation

Following the identification of patients prescribed anticoagulants, the potential costs associated with the transition to the newer agents were calculated. Cost-effectiveness and benefit-to-risk profiles are a common method for determining the resource costs incurred in achieving a predefined clinical outcome. While several perspectives may be adopted, this evaluation has considered the view of the third-party payer (the Pharmaceutical Benefits Scheme [PBS]) and has calculated only direct costs.

The cost of year of analysis was 2009. The costs of giving dabigatran or rivaroxaban to prevent VTE and SPAF assumed the mean duration of prophylaxis in VTE and SPAF under the Australian Therapeutics Guidelines (2009). Drug costs were taken from the Australian Pharmaceutical Benefits Scheme, July 2009. Additionally, the cost for rivaroxaban included a 33% supplier-rebate based on a pricing submission to the hospital for its supply. Time for the administration of warfarin included 20 minutes of nurse time to check INRs and to administer the drug. The administration costs for enoxaparin included time to administer injection and to counsel outpatients on self-injection following discharge. The associated costs were calculated from the nurse hourly rates from Australian public health data (2007-2008). Administration costs for dabigatran and rivaroxaban included 10 minutes of nursing time to retrieve the dose of medication.

The anticoagulant effect of dabigatran can be measured by ecarin clotting time and thrombin time. Similarly, rivaroxaban may be monitored by measuring factor Xa activity as well as prothrombin time. However, such tests have not been widely used in association with dabigatran and rivaroxaban therapy. While they are potentially advantageous, regular testing of alanine aminotransferase (ALT) levels has been recommended at baseline and monthly for the first 6 months of commencing therapy, every second month for the remainder of the first year, and every third month thereafter, as a safety precaution. Given that all patients admitted to hospital will receive a liver function test which measures ALT as part of their standard assessment, only the cost of the second test was included as part of long-term monitoring and follow-up. The cost of conducting ALT testing was determined accordingly and derived from the Medicare Benefits Schedule, July 2009.

Monitoring costs in conducting INR tests for warfarin patients was derived from the Medicare Benefits Schedule July 2009. No monitoring tests were included for enoxaparin since the dose is calculated according to the specific weight and renal function of each individual patient, as recommended by the Australian Therapeutic Guidelines (2009).

The length of stay for confirmed cases of VTE and atrial fibrillation (AF) was modelled for data from the prospective audit. A weighted average of the incremental mean length of stay was determined (8.5 days for VTE, 12.5 days for AF). The cost associated with the length of stay was estimated using a per diem cost for each ward. The cost of a hospital admission due to a major bleeding event was based on 2007-2008 Australian public hospital data. Clinically relevant minor bleeding events were estimated as two outpatient visits and were based on clinical opinion, as has been previously described.

Analysis

Given that the basic analysis adopts a modelling approach, it was sensible to apply a sensitivity analysis to judge how robust the results were based on assumptions made. The following data were subjected to sensitivity analysis: (1) cost of monitoring anticoagulation; (2) length of stay for bleeding events; (3) percentage of patients admitted to the hospital for a bleeding event; (4) percentage of patients discharged to hospital continuing care; (5) unit costs for hospital bed-days; and (6) incidence of bleeding events in NSW.

A cost minimisation analysis was applied to determine cost differences between each anticoagulant therapy given that the incidence of VTE/SPAF and bleeding events were equivalent in each treatment group.

RESULTS

In total, over the 2-week clinical audit, 67 patients were identified from the orthopaedic and coronary care units who had VTE or SPAF and were administered warfarin 5mg once daily (n=46) or enoxaparin 40mg intravenously once daily (n=21) (the approved doses in patients for the prevention of VTE or SPAF in Australia). Baseline characteristics were well matched between the treatment groups for each traditional anticoagulant (Table 1).
Intracranial Surgical Site Gastrointestinal Other
No. of Bleeding Events 1 (1.82%) 2 (3.62%) 3 (5.45%) 2 (3.64%)
Survivor Permanently Disabled 1 (1.82%) 0 0 0
Cost (AUS) Acute Care [28] 2420 4840 7260 4840
Long-term Care [29] 8712 0 0 0

* Data are number (%) or mean (SD). +Body-mass index is the weight in kilograms divided by the square of the height in metres.
Braidy N, Bui K, Bajorek B. Evaluating the impact of new anticoagulants in the hospital setting. Pharmacy Practice (Granada) 2011 Jan-Mar;9(1):1-10.

Figure 3. Incidence of major and minor bleeding events with warfarin and enoxaparin

Table 3. Cost of prophylaxis for VTE/SPAF for patients on warfarin, enoxaparin, dabigatran or rivaroxaban

|                         | Warfarin 5mg Once Daily (n=39) | Enoxaparin 40mg Once Daily (n=16) | Dabigatran 220mg Once Daily | Rivaroxaban 10mg Once Daily |
|-------------------------|---------------------------------|-----------------------------------|-----------------------------|-----------------------------|
| Cost of Prophylaxis     | VTE (AUD)                       | SPAF (AUD)                        | VTE (AUD)                   | SPAF (AUD)                   |
| No. of doses received   | 10.44 (1.9)                     | 9.66 (1.5)                        | 16.01 (2.1)                 | 0 (1.3)                     |
| during admission (SD)   |                                 |                                  |                             |                             |
| Cost of drug per day    | 5.85                            | 5.85                              | 6.12                        | 6.12                        |
| (AUD)                   |                                 |                                  |                             |                             |
| Administration Costs    |                                 |                                  |                             |                             |
| per day (AUD)           | Inpatient 10.35                  | 15.40                             | 5.18                        | 5.18                        |
|                         | Outpatient 0                     | 0                                 | 5.44                        | 0 (5.18)                    |
| Cost per patient (AUD)  | 71.42                           | 118.86                            | 73.49                       | 113.39                      |
|                         |                                 |                                  |                             |                             |
|                         | Administration costs were calculated as 20 minutes per patient from the nurse hourly rates from Australian public health data (2007-2008).|

Table 4. Cost of monitoring for VTE/SPAF for patients on warfarin, enoxaparin, dabigatran or rivaroxaban

|             | Warfarin 5mg Once Daily (n=39) | Dabigatran 220mg Once Daily | Rivaroxaban 10mg Once Daily |
|-------------|---------------------------------|-----------------------------|-----------------------------|
| Cost of Monitoring | INR (95% CI)                           | ALT (95% CI)               | ALT (95% CI)               |
| No. of tests conducted | 3.23 (1.44)                          | 3.71 (1.59)                | 2                           | 2                            |
| during admission (SD)   |                                 |                             |                             |                             |
| Cost of test per day (AUD) | 13.80                              | 9.75                        | 9.75                        | 9.75                        |
| Cost per patient (AUD)  | 44.57                            | 9.75                        | 9.75                        | 9.75                        |

* The initial ALT test was not included in the cost analysis since the it is conducted as routine in every patient that is admitted.

Table 5. Efficacy of warfarin, enoxaparin, dabigatran or rivaroxaban for the prophylaxis of VTE and SPAF

|                         | RR, Dabigatran vs Warfarin (95% CI) | RR, Dabigatran vs Enoxaparin (95% CI) | RR, Rivaroxaban vs Warfarin (95% CI) | RR, Rivaroxaban vs Enoxaparin (95% CI) |
|-------------------------|------------------------------------|--------------------------------------|--------------------------------------|--------------------------------------|
| Efficacy                | VTE Stroke                         | VTE Stroke                            | VTE Stroke                            | VTE Stroke                            |
| Total VTE/stroke and all-cause mortality | 0.94 (0.76-1.39)                  | 0.88 (0.58-1.39)                     | 0.97 (0.83-1.13)                     | 0.71 (0.55-1.05)                     |
| RR (95% CI)             | VTE Stroke                         | VTE Stroke                            | VTE Stroke                            | VTE Stroke                            |
| Total VTE/stroke and all-cause mortality | 0.68 (0.40-0.66)                  | 0.68 (0.31-0.75)                     | 0.68 (0.40-0.66)                     | 0.68 (0.31-0.75)                     |
| Incidence (% n/N)       | 37.7 (193/51)                      | 3.9 (87/2331)                        |                                     |                                     |
Table 6. Incidence of bleeding events for warfarin, enoxaparin, dabigatran or rivaroxaban for the prophylaxis of VTE and SPAF

|                  | RR, Dabigatran vs Warfarin (95% CI) | RR, Dabigatran vs Enoxaparin (95% CI) | RR, Rivaroxaban vs Warfarin (95% CI) | RR, Rivaroxaban vs Enoxaparin (95% CI) | Incidence (% n/N) (95%CI) |
|------------------|------------------------------------|---------------------------------------|-------------------------------------|---------------------------------------|--------------------------|
|                  | VTE SPAF                           | VTE SPAF                              | VTE SPAF                            | VTE SPAF                              |                          |
| Major bleeding   |                                    |                                       |                                     |                                       |                          |
| Event            | 1.69 (0.81-3.52)                   | 1.35 (0.46-2.78)                      | 1.33 (0.71-2.18)                    | 1.14 (0.71-1.82)                      | 1.3 (9694)               |
| Minor bleeding   |                                    |                                       |                                     |                                       |                          |
| Event            | 1.86 (0.64-2.26)                   | 1.04 (0.62-1.33)                      | 1.41 (0.75-2.27)                    | 1.12 (0.81-1.86)                      | 15.3 (106/694)          |

Figure 4. Total cost of anticoagulation therapy over total period of hospitalisation per patient for VTE and SPAF

Table 7. Cost-effectiveness analysis of dabigatran and rivaroxaban versus warfarin and enoxaparin in VTE and SPAF

| Cost Difference              | Dabigatran Dominant | Dabigatran Dominant | Rivaroxaban Dominant | Rivaroxaban Dominant |
|------------------------------|---------------------|---------------------|----------------------|----------------------|
| VTE                          |                     |                     |                      |                      |
| SPAF                         |                     |                     |                      |                      |
| Dabigatran Dominant          |                     |                     |                      |                      |
| Rivaroxaban Dominant         |                     |                      |                      |                      |

If new anticoagulants are to be considered as potential replacements for warfarin and/or enoxaparin for preventing VTE’s or stroke associated with atrial fibrillation, drug cost is a critical issue. To determine the potential cost benefits for dabigatran and rivaroxaban compared to traditional anticoagulants, the cost of using these newer anticoagulants was compared to the cost of warfarin and enoxaparin for VTE and SPAF respectively (Table 3). The costs of warfarin (5mg once daily) and enoxaparin (40mg once daily) for these indications is AUD5.85 and AUD6.12 per day, respectively. The cost of dabigatran (220mg once daily) was significantly higher compared to traditional anticoagulants at AUD8.87. The cost of rivaroxaban (10mg once daily) was significantly lower than for the traditional anticoagulants, at AUD4.27 after a 33% rebate under a supplier agreement. This represents the best available estimates for the costs of using dabigatran and rivaroxaban, although the dose might vary for VTE and SPAF depending on patient’s renal and hepatic function. The cost of prophylaxis was significantly higher for warfarin and enoxaparin compared to dabigatran and rivaroxaban (Table 3). The costs for enoxaparin were higher due to the cost of nursing time for administration of subcutaneous injections during inpatient stay and the cost of administration after discharge. Administration costs included the cost of nursing time for frequent and accurate monitoring of INR levels in inpatients and dosage retrieval for dabigatran and rivaroxaban. The reduced cost of dabigatran was due to repackaging the half-dose given on the first day of prophylaxis. The cost of rivaroxaban was significantly lower than warfarin and enoxaparin, primarily due to supplier rebates.

Routine monitoring of warfarin (i.e. international normalised ratio -INR) is a standard guideline for the prophylaxis of VTE and SPAF. The cost of monitoring anticoagulation was significantly higher for warfarin (AUD44.57 and AUD51.20 respectively) compared to dabigatran (AUD9.75) and rivaroxaban (AUD9.75) (Table 4). While the cost per test for INR is only slightly higher than the cost of ALT test, the cost difference is attributed to the greater frequency of monitoring of INR levels compared to ALT testing (Table 4). No monitoring costs were included for enoxaparin.

The results for the efficacy of anticoagulation are presented in Table 5. The outcomes were comparable for patients receiving warfarin, enoxaparin, dabigatran or rivaroxaban in both the VTE and SPAF analysis. Although the incidence of VTE/SPAF was slightly lower for dabigatran and rivaroxaban compared to warfarin and enoxaparin, the differences in efficacy were small (Table 5). Although the incidence of bleeding events (including major and minor bleeds) was slightly higher for dabigatran and rivaroxaban compared to warfarin and enoxaparin, it did not reach statistical significance (Table 6). Altogether, these results reflect a similar efficacy and safety profile as for the new anticoagulants in the RE-MODEL/RE-NOVATE and RECORD/PETRO trials, respectively.
In patients undergoing prophylaxis for VTE and SPAF, dabigatran was dominant over warfarin and enoxaparin in terms of cost of drug administration and therapeutic monitoring, and efficacy and safety of the anticoagulant therapy (Figure 4 and Table 7). Similarly, rivaroxaban was dominant over warfarin and enoxaparin for the prophylaxis of VTE and SPAF. While the cost of therapeutic monitoring is significantly reduced for rivaroxaban compared to warfarin, the presence of existing supplier-rebates further reduces the costs of using this newer agent, compared to the costs of warfarin and enoxaparin which maintain similar efficacy and safety profiles.

**DISCUSSION**

VTE and stroke are extremely common and often fatal conditions, yet there are poorly managed due to limited treatment option. Not surprisingly, this has led to the development of newer anticoagulants. Traditionally, warfarin has been used in the hospital setting for the prophylaxis of VTE and SPAF. However, it has several other shortcomings, including: (1) the need for frequent therapeutic monitoring, which is a considerable inconvenience to the patient and a significant cost burden on the healthcare system; (2) INR levels are significantly affected by foods and herbs, adding to the difficulty of maintaining dosage regimens; (3) apart from GI bleeding, warfarin may also cause haemorrhagic stroke, osteoporosis and bone fracture (long-term use) and other debilitating complications; and (4) warfarin can interact with at least 60 commonly administered drugs including paracetamol and can be as fatal as elevating INR to 6.0 or higher. LMWH’s such as enoxaparin are also effective at preventing coagulation but are less commonly used due to the need for administration by injection. Conceived efforts to find alternative anticoagulants have led to the development of the new oral anticoagulants, dabigatran and rivaroxaban.

Dabigatran and rivaroxaban have been previously reported to have comparable efficacy and similar safety profiles compared to warfarin and enoxaparin. The current study results indicate that dabigatran may be cost effective compared to warfarin and enoxaparin for use in patients undergoing prophylaxis for VTE and SPAF, respectively. In VTE and SPAF patients, cost parameters and outcomes were similar for these anticoagulants. Modest cost savings were expected for dabigatran for the prevention of VTE since it can provide oral anticoagulant therapy without the need for dose adjustment or routine coagulation monitoring. In SPAF patients given warfarin, and in whom long-term prophylaxis is necessary, estimated cost savings are more substantial. Moreover, monitoring in some countries is by the physician or pharmacist and warfarin monitoring costs would therefore be higher than the costs presented in this study. These findings are consistent with a similar study by the United Kingdom (UK) Health Service, whereby dabigatran was found to be cost-effective compared with the dose of enoxaparin approved for use in patients undergoing orthopaedic surgery in the UK.

Moreover, the longer half-life of dabigatran makes it an ideal agent for stroke prevention. The RE-LY (Randomized Evaluation of Long-Term Anticoagulation Therapy) evaluated the effect of two doses of dabigatran (150 and 110 mg) for SPAF relative to warfarin. While higher doses were superior to warfarin for stroke prevention, similar efficacy to warfarin was observed at lower doses. Although no difference in bleeding rates were reported for warfarin compared to higher doses of dabigatran, lower doses were associated with lower incidence of bleeding events. This may further direct the cost-efficacy results in the favour of dabigatran.

Similarly, rivaroxaban was found to be more cost-effective compared to warfarin and enoxaparin in the current study. On the basis of the pharmacology and the findings of this study and others, treatment with rivaroxaban might become a useful adjunct to therapy in patients for the prophylaxis of VTE and/or SPAF. As previously mentioned, its oral route of administration, low pharmacological dose, and timing of administration are more advantageous compared to warfarin and enoxaparin. Moreover, under a plausible set of assumptions, the reduced cost of rivaroxaban make it more cost effective compared to traditional anticoagulant therapy.

Several other phase III trials, such as MAGELLAN and EINSTEIN are currently examining the impact of rivaroxaban in other clinical settings such as pulmonary embolism. If rivaroxaban is found to be more efficacious than traditional anticoagulants, then it may replace warfarin and/or enoxaparin for initial and acute therapy as well as for long-term care. The ROCKET study is aimed at comparing a fixed dose of rivaroxaban in AF and SPAF compared to standard warfarin treatment. ATLAS TMI 5 is also evaluating the effectiveness of rivaroxaban for the treatment of acute coronary syndrome. Results of both studies are expected to be released in 2011. Other factor Xa inhibitors have been developed for the prevention of VTE, for which warfarin, has long been the first choice. Apixaban is being tested to prevent DVT and pulmonary embolism that sometimes occur within patients hospitalized for acute medical illness (ADOPT). These studies are almost complete and will likely to be published in the coming months. The use of new oral anticoagulants poses several benefits over warfarin and enoxaparin for the prophylaxis of VTE and stroke. Firstly, the oral route of administration significantly reduces the need for self-administration in outpatients prescribed enoxaparin, or the need for further healthcare requirements in patients unable to self-inject following hospital discharge.

These issues limit the use of enoxaparin for extended prophylaxis. Other costs that may be reduced include the cost of platelet monitoring for HITs, needle stick injuries and disposal of sharps. Newer oral anticoagulants may also improve patient compliance due to reduced need for frequent monitoring and dosage adjustments, or may prefer...
oral medications rather than subcutaneous injections.39

In the present study, modelling was necessary to estimate the incidence and impact of VTE/SPAF and bleeding events. While the model parameters were validated by feedback from clinical experts, like all models, assumptions about these may affect the final results. For instance, when a single patient experienced more than one event during their hospital admission, (e.g. VTE and/or stroke, and bleeding events), the costs of managing the patient were assumed to be additive. Moreover, the incidence of VTE/SPAF from the descriptive data indicated that the probability of having at least one event during the study period and does not reflect the possibility of multiple events occurring in an individual (e.g VTE and stroke). In addition, the current study does not include additional costs associated with platelet monitoring for heparin induced thrombosis (HITs), possible needle-stick injuries or the disposal of sharps. The study does not include potential utility costs such as avoidance of daily enoxaparin injection. Therefore, the cost effectiveness for dabigatran and rivaroxaban may be underestimated when compared to enoxaparin.39

Furthermore, the current study was carried out over a very short two-week period based on the perspective of Ryde Community Hospital, which is a community teaching hospital with limited number of beds. Therefore, the number of patients able to be included in the warfarin/enoxaparin arm is small. Therefore, there are limitations with applying the drug costs involved here to other healthcare facilities operating under different contracts or health systems internationally, and where the number of patients hospitalised may be much greater than in the present study.

While the cost of life is not included in the current analysis, the cost efficacy may likely to be improved, given the lower mortality of dabigatran and rivaroxaban over warfarin and enoxaparin for the treatment of VTE/SPAF. Using data collected from the RE-LY trial and other published studies of antiocoagulation, Freeman et al (2010) recently showed that dabigatran can improve the quality of life of patients with SPAF compared to traditional adjusted-dose warfarin in the United Kingdom.46 Similarly, rivaroxaban was shown to be more cost-effective compared to enoxaparin for the treatment of VTE in the Irish hospital setting. It remains uncertain whether dabigatran is more cost effective than rivaroxaban when compared to warfarin or enoxaparin, using the same strategy.47

CONCLUSION
In conclusion, this study highlights the potential cost-effectiveness of newer anticoagulants, dabigatran and rivaroxaban, compared to warfarin and enoxaparin. These agents may offer economic advantages, as well as clinical benefits, in the hospital-based management of anticoagulated patients.

CONFLICT OF INTEREST
The authors have no conflicts of interest to declare.

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