Coronary and mortality risk of novel oral antithrombotic agents: a meta-analysis of large randomised trials

Koon-Hou Mak

ABSTRACT
Objective: Oral direct thrombin and anti-Xa inhibitors have been shown to be efficacious in the prevention and treatment of venous thromboembolism, and prevention of embolic events in atrial fibrillation. Recent studies showed that dabigatran may be associated with increased rates of myocardial infarction (MI). Coronary risk for the other agents was unclear. The aim of the study is to determine the coronary risk among four novel antithrombotic agents.

Design: Mixed treatment comparison meta-analysis.

Data sources and study selection: Randomised controlled trials (RCTs) on ximelagatran, dabigatran, rivaroxaban and apixaban were obtained from PubMed search (February 2012) and major scientific meeting in 2011. The random-effects model was used to evaluate the effect of these agents on MI or acute coronary syndrome (MI/ACS), major bleeding complication and all-cause mortality.

Results: From 28 RCTs (n=138 948), the risk for MI/ACS was higher for dabigatran (OR 1.30; 95% CI 1.04 to 1.63; p=0.021) but lower for rivaroxaban (OR 0.78; 95% CI 0.69 to 0.89; p<0.001). Ximelagatran showed a higher risk for MI/ACS, which was not statistically significant, while apixaban demonstrated a non-significant lower likelihood. Among the RCTs for MI/ACS among the four agents, only those pertaining to ximelagatran showed heterogeneity. Major bleeding complication rates varied considerably among different agents. Importantly, these agents were associated with a lower all-cause mortality, without heterogeneity among the studies.

Conclusions: The risk for coronary events was significantly higher for dabigatran but not significantly higher for ximelagatran. Conversely, this risk was lower among anti-Xa inhibitors. All-cause mortality was lower among those receiving novel antithrombotic agents. This information may be useful in selecting agents for specific subsets of patients requiring anticoagulation.

INTRODUCTION
Several cardiovascular conditions are related to thromboembolism. In the past few decades, focus has been on the development of antiplatelet agents because of the perceived pre-eminent role played by thrombocytes in arterial, particularly coronary, thrombosis. For several years, advancement in anticoagulation has been limited to refining the heparin complex and parenteral direct thrombin inhibitors such as hirudin and bivalrudin. Indeed, warfarin was the sole oral anticoagulant for the past 60 years. Novel agents have been designed to act against factor Xa and thrombin recently. Their efficacy has been shown in preventing venous thromboembolism (VTE) among patients undergoing hip or knee surgery and embolic events among those with atrial fibrillation, and treating those with VTE or acute coronary syndromes (ACS). Amidst the enthusiasm of favourable results, higher rates of myocardial infarction (MI) among patients receiving dabigatran initially reported in the Randomised Evaluation of Long-term Anticoagulant Therapy (RE-LY) trial have generated concern regarding the overall
Oral novel antithrombotic agents and coronary risk

effectiveness of this agent. Although subsequent re-analysis of the data following identification of another four clinical and 28 silent MI showed that the increase was not statistically significant (HR 1.28; 95% CI 0.98 to 1.67; p=0.07). However, a recent meta-analysis showed that the risk of coronary events was increased with the use of dabigatran, even after including the additional events. To date, there have been four novel oral anticoagulants that have been evaluated in large clinical trials for thromboembolic conditions: ximelagatran (Exanta; Astra-Zeneca, London, UK), dabigatran (Pradaxa; Boehringer Ingelheim, Ingelheim, Germany), rivaroxaban (Xarelto; Bayer, Leverkusen, Germany) and apixaban (Eliquis; Bristol-Myers Squibb, New York City, New York, USA). The first two agents are direct thrombin inhibitors while the latter two act against factor Xa. With different mechanism of action, the aim of the study is to review the risk of acute coronary events among these agents. Taken together, they may provide a better ascertainment of the coronary risk between direct thrombin inhibitors and anti-Xa agents.

RESULTS

A total of 274 abstracts were identified and reviewed. Of these, 42 full-text articles were appraised, and eventually, 28 randomised control trials (RCTs) were selected (figure 1), consisting of 138 948 participants. The numbers of trials evaluating ximelagatran, dabigatran, rivaroxaban and apixaban were six, nine, seven and seven, respectively, and were sponsored by their respective pharmaceutical companies. They were conducted in the setting VTE prevention among patients undergoing hip or knee surgery (13 studies), treatment of individuals with VTE (5 studies), prevention of embolic events in patients with atrial fibrillation (6 studies) and treatment of subjects with ACSs (4 studies). Study participants were followed from about a week to 2 years. The characteristics of the trials are provided in table 1.

Impact on MI/ACS

Of the four drugs, the risk for MI/ACS was higher for dabigatran (OR 1.30; 95% CI 1.04 to 1.63; p=0.021) but lower for rivaroxaban (OR 0.78; 95% CI 0.69 to 0.89; p<0.001) (figure 2). The other oral direct thrombin inhibitor, ximelagatran, showed a higher risk for MI/ACS, which was not statistically significant, and apixaban, a factor Xa inhibitor, demonstrated a non-statistically significant lower likelihood. Unlike trials involving dabigatran, rivaroxaban and apixaban, there was marked heterogeneity for studies evaluating ximelagatran (I²=79.69; p=0.007).

Major bleeding complications

Overall, the risk of major bleeding complications was comparable between oral direct thrombin inhibitors and warfarin (figure 3). When the trial on patients with ACS was excluded, dabigatran was associated with a reduced risk for major bleeding complications (OR 0.89; 95% CI 8.80 to 0.999; p=0.049). But there was still considerable heterogeneity among the studies (I²=67.29; p=0.003). Conversely, the risk for major bleeding complication was 15% higher for rivaroxaban. Again, there was marked heterogeneity because of dissimilar trial design, the heightened risk for major bleeding complication was attenuated after excluding the study on ACS (OR 1.03;
95% CI 0.90 to 1.19; p=0.638). Test for heterogeneity became non-significant ($I^2=3.32$; $p=0.395$). Overall, apixaban was associated with a non-statistically significant lower likelihood for major bleeding, with marked heterogeneity among trials. When the studies on ACS were excluded, the risk for major bleeding of significantly lower for apixaban (OR 0.69; 95% CI 0.61 to 0.79; $p<0.001$) and without significant heterogeneity ($I^2=1.84$). Conversely, major bleeding complications occurred more frequently among patients receiving apixaban in ACS trials (OR 2.61; 95% CI 1.52 to 4.72; $p<0.001$) without significant heterogeneity ($I^2<0.001$).

**All-cause mortality**

Aside from ximelagatran, the use of dabigatran, rivaroxaban and apixaban was associated with the reduction in all-cause mortality (figure 4). Importantly, there was no significant heterogeneity among the trials.

Funnel plot with Engger regression test did not show evidence for publication bias for the various outcomes (figure 5 showing data only for MI/ACS). Meta-regression analysis did not show any relationship between each antithrombotic agent and individual outcome measures, except for ximelagatran with MI/ACS ($p=0.007$), rivaroxaban with major bleeding complication ($p<0.001$) and apixaban with major bleeding complication ($p=0.004$).

**DISCUSSION**

This meta-analysis showed that dabigatran was associated with increased risk for acute coronary events. Conversely, the greater likelihood for coronary events for the other oral direct thrombin inhibitor, ximelagatran, was not statistically significant. The excess risk associated with dabigatran was comparable to the findings of the earlier meta-analysis. Conversely, the risk for MI/ACS was lower for rivaroxaban, and a non-statistically significant reduction was observed for apixaban. Therefore, it appeared that the coronary risk differed between oral direct thrombin inhibitors and anti-Xa agents. Although the variation in the use of antiplatelet agents could have accounted for some of these differences, it was interesting to note that dabigatran was associated with a higher and rivaroxaban was associated with a lower risk for MI/ACS in clinical studies of ACS patients. Majority of them would have been treated with at least one antiplatelet agent. Therefore, based on these findings, those with heightened coronary risk, the use of anti-Xa agents may be preferable to direct thrombin inhibitors.

While both ximelagatran and low-molecular-weight heparins were able to reduce platelet activation, thrombin generation and endogenous thrombin potential, the time reduction for endogenous thrombin potential was greater for dalteparin compared with ximelagatran. Conversely, rivaroxaban was superior to dalteparin in preventing thrombin generation following hip
<table>
<thead>
<tr>
<th>Study name</th>
<th>Study population</th>
<th>Primary endpoint</th>
<th>Coronary event</th>
<th>Study drug and dose (number of subjects)</th>
<th>Control drug and dose (number of subjects)</th>
<th>Duration of therapy</th>
<th>Jadad score</th>
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<tr>
<td>Venous thromboembolism prophylaxis</td>
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<td>EXULT A⁷</td>
<td>Knee surgery</td>
<td>VTE, death</td>
<td>NR</td>
<td>Ximelagatran 36 mg twice daily (n=629)</td>
<td>Warfarin (n=608)</td>
<td>7–12 days</td>
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<td>Knee surgery</td>
<td>VTE, death</td>
<td>NR</td>
<td>Ximelagatran 36 mg twice daily (n=982)</td>
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<td>Hip surgery</td>
<td>VTE, death</td>
<td>ACS</td>
<td>Dabigatran 150 mg once daily (n=874)</td>
<td>Enoxaparin 40 mg once daily (n=897)</td>
<td>28–35 days</td>
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<td>Knee surgery</td>
<td>VTE, death</td>
<td>ACS</td>
<td>Dabigatran 150 mg once daily (n=526)</td>
<td>Enoxaparin 40 mg once daily (n=512)</td>
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<td>RE-MOBILIZE¹¹</td>
<td>Knee surgery</td>
<td>VTE, death</td>
<td>cardiac events*</td>
<td>Dabigatran 150 mg once daily (n=649)</td>
<td>Enoxaparin 30 mg twice daily (n=643)</td>
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<td>MI</td>
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<td>Hip surgery</td>
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<td>MI</td>
<td>Rivaroxaban 10 mg daily (n=1595)</td>
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<td>36† (30–42) days</td>
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<td>MI</td>
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<td>MI</td>
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<td>MI</td>
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<td>ADVANCE 1¹⁷</td>
<td>Knee surgery</td>
<td>VTE, death</td>
<td>MI</td>
<td>Apixaban 2.5 mg twice daily (n=1599)</td>
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<td>VTE, death</td>
<td>MI</td>
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<td>Hip surgery</td>
<td>VTE, death</td>
<td>MI</td>
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**Treatment of venous thromboembolism**

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<th>Study name</th>
<th>Study population</th>
<th>Primary endpoint</th>
<th>Coronary event</th>
<th>Study drug and dose (number of subjects)</th>
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<td>Recurrent VTE</td>
<td>ACS</td>
<td>Ximelagatran 36 mg twice daily (n=1240)</td>
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<td>VTE therapy</td>
<td>VTE</td>
<td>ACS</td>
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<td>RE-SONATE²²</td>
<td>Extended VTE therapy</td>
<td>Recurrent VTE, related death</td>
<td>CV events</td>
<td>Dabigatran 150 mg twice daily (n=681)</td>
<td>Warfarin (n=1426)</td>
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<td>REMEDY²³</td>
<td>Extended VTE therapy</td>
<td>Recurrent VTE, related death</td>
<td>ACS</td>
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<td>Symptomatic DVT therapy</td>
<td>Recurrent VTE</td>
<td>ACS</td>
<td>Rivaroxaban 15 mg twice daily for 3 weeks, then 20 mg daily (n=1718)</td>
<td>Heparin followed by warfarin (n=1711)</td>
<td>3, 6, 12 months</td>
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<th>Study name</th>
<th>Study population</th>
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<td>Stroke and embolic events MI</td>
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<td>Ximelagatran 36 mg twice daily (n=1704)</td>
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<td>SPORTIF V</td>
<td>Non-valvular atrial fibrillation</td>
<td>Stroke and embolic events MI</td>
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<td>Ximelagatran 36 mg twice daily (n=1960)</td>
<td>Warfarin (n=1962)</td>
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<td>Warfarin (n=6022)</td>
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<td>Non-valvular atrial fibrillation</td>
<td>Stroke or embolic events MI</td>
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<td>Warfarin (n=7004)</td>
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<td>Atrial fibrillation warfarin unsuitable</td>
<td>Stroke or embolic events MI</td>
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<td>Apixaban 5/2.5 mg twice daily (n=2808)</td>
<td>Aspirin 81–324 mg daily (n=2791)</td>
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<td>Stroke or embolic events MI</td>
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<td>Apixaban 5/2.5 mg twice daily (n=9120)</td>
<td>Warfarin (n=9081)</td>
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<td>RE-DEEM</td>
<td>STE or NSTE MI</td>
<td>CV death, MI, stroke</td>
<td>ACS</td>
<td>Dabigatran 50 mg twice daily (n=369)</td>
<td>Placebo (n=371)</td>
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<td>75 mg twice daily (n=368)</td>
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<td>110 mg twice daily (n=406)</td>
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<td>150 mg twice daily (n=347)</td>
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<td>ATLAS ACS 2-TIMI 51</td>
<td>Unstable angina, STE or NSTE MI</td>
<td>CV death, MI, stroke</td>
<td>CV death or MI</td>
<td>Rivaroxaban 2.5/5 mg twice daily (n=10229)</td>
<td>Placebo (n=5113)</td>
<td>13 months§</td>
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<td>APPRAISE</td>
<td>Unstable angina, STE or NSTE MI</td>
<td>CV death, MI, re-ischemia or ischemic stroke</td>
<td>CV death or MI</td>
<td>Apixaban 10 mg daily (n=315)</td>
<td>Placebo (n=599)</td>
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<td>2.5 mg twice daily (n=315)</td>
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<td>APPRAISE 2</td>
<td>Unstable angina, STE or NSTE MI</td>
<td>CV death, MI, stroke</td>
<td>ACS</td>
<td>Apixaban 5 mg twice daily (n=3705)</td>
<td>Placebo (n=3687)</td>
<td>240 days†</td>
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</table>

*Cardiac events, specifics were not provided but events were reviewed by a blinded independent committee.
†Median.
‡Limited information available.
§Mean.
VTE, venous thromboembolism; NR, not reported; ACS, acute coronary syndrome (consisting of unstable angina, myocardial infarction and cardiac death); MI, myocardial infarction; CV, cardiovascular; STE, ST-segment-elevation; NSTE, non-ST-segment-elevation

Acronyms for studies, where applicable: EXULT, Exanta Used to Lessen Thrombosis; RECORD, Regulation of Coagulation in Orthopedic Surgery to Prevent Deep Vein Thrombosis and Pulmonary Embolism; ADVANCE, Apixaban Dose Orally versus. Anticoagulant with Enoxaparin; THRIVE, the Thrombin Inhibitor in Venous Thromboembolism; SPORTIF, Stroke Prevention using an Oral Thrombin Inhibitor in atrial Fibrillation; RE-LY, Randomized Evaluation of Long-Term Anticoagulation Therapy; ROCKET AF: Rivaroxaban Once Daily Oral Direct Factor Xa Inhibition Compared with Vitamin K Antagonism for Prevention of Stroke and Embolism Trial in Atrial Fibrillation; AVERROES, Apixaban Versus Acetylsalicylic Acid to Prevent Stroke in Atrial Fibrillation Patients Who Have Failed or Are Unsuitable for Vitamin K Antagonist Treatment; ARISTOTLE, Apixaban for Reduction in Stroke and Other Thromboembolic Events in Atrial Fibrillation; ATLAS ACS 2-TIMI 51, Anti-Xa Therapy to Lower Cardiovascular Events in Addition to Standard Therapy in Subjects with Acute Coronary Syndrome-Thrombolysis in Myocardial Infarction 51; APPRAISE, Apixaban for Prevention of Acute Ischemic and Safety Events Trial.
and knee replacement surgery\textsuperscript{37} and reduces tissue factor induced platelet aggregation.\textsuperscript{38} In vitro studies indicated that direct thrombin inhibitors were associated with paradoxical coagulation compared with factor Xa inhibitors,\textsuperscript{39} which was likely mediated by preventing thrombin-induced activation of Protein C. This is a

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{Figure2.png}
\caption{Risk of coronary events.}
\end{figure}

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{Figure3.png}
\caption{Risk for major bleeding complications.}
\end{figure}

with paradoxical coagulation compared with factor Xa inhibitors,\textsuperscript{39} which was likely mediated by preventing thrombin-induced activation of Protein C. This is a
natural anticoagulant and part of the negative feedback system after thrombin generation. Furthermore, inflammatory markers were increased with long-term use of direct oral thrombin inhibitors. Urinary 11-dehydrothromboxane \( \beta_2 \) was elevated for those receiving dabigatran compared with warfarin among 502 patients.

Figure 4 Risk for all-cause mortality.

Figure 5 Fixed-effects funnel plot with Egger regression test for the evaluation of publication bias for coronary events.
<table>
<thead>
<tr>
<th>Study name</th>
<th>Major bleeding complication definition</th>
<th>Antiplatelet agent</th>
</tr>
</thead>
</table>
| Venous thromboembolism prophylaxis | **EXULT A**<sup>7</sup> Occurrence of at least one of the following:  
1. Critical site (intracranial, retroperitoneal, intraocular, intraspinal, pericardial)  
2. Bleeding index $\geq 2.0$ (difference between baseline and postbleeding haemoglobin level (g/l) plus number of packed cells or whole blood transfusion)  
3. Need for medical or surgical intervention at operative site  
4. Fatal | Not allowed |
|                            | **EXULT B**<sup>8</sup> Not clearly stated                                                                                                                                                                                                 | Not allowed |
|                            | **RE-NOVATE**<sup>9</sup> Acute overt clinical bleeding with one of the following:  
1. Critical site (intracranial, retroperitoneal, intraocular, intraspinal, pericardial)  
2. Fall in haemoglobin $\geq 20$ g/l in excess of that expected by investigator  
3. Transfusion $\geq 2$ units of packed cells or whole blood in excess of that expected by investigator  
4. Leading to re-operation  
5. Warranting treatment cessation  
6. Fatal | Aspirin dose $<162$ mg daily permitted |
|                            | **RE-MODEL**<sup>10</sup> As in RE-NOVATE                                                                                                                                                                                               | Aspirin dose $<160$ mg daily permitted |
|                            | **RE-MOBILIZE**<sup>11</sup> Occurrence of at least one of the following:  
1. Symptomatic intracranial, retroperitoneal, intraocular or intraspinal bleeding  
2. Clinically overt bleeding with fall of haemoglobin $\geq 2.0$ g/dl and/or leading to transfusion of $\geq 2$ units of packed cells or whole blood  
3. Need for treatment cessation or surgical intervention at operative site  
4. Fatal | Aspirin dose $<160$ mg daily permitted |
|                            | **RE-NOVATE II**<sup>12</sup> As in RE-NOVATE                                                                                                                                                                                              | Aspirin dose $<162$ mg daily permitted |
|                            | **RECORD**<sup>13</sup> Occurrence of at least one of the following:  
1. Intracranial, retroperitoneal, intraocular or intraspinal bleeding  
2. Clinically overt bleeding with fall of haemoglobin $\geq 2.0$ g/dl  
3. Transfusion of $\geq 2$ units of packed cells or whole blood  
4. Need for surgical intervention at operative or bleeding site  
5. Fatal | | |
|                            | **RECORD2**<sup>14</sup> Occurrence of at least one of the following:  
1. Critical site bleeding; for example, intracranial, retroperitoneal, intraocular or intraspinal  
2. Clinically overt bleeding with fall of haemoglobin $\geq 2.0$ g/dl (calculated from first post-operative level)  
3. Transfusion of $\geq 2$ units of packed cells or whole blood  
4. Need for surgical intervention at operative or bleeding site  
5. Fatal | | |
|                            | **RECORD3**<sup>15</sup> Occurrence of at least one of the following:  
1. Critical organ bleeding  
2. Clinically overt bleeding with fall of haemoglobin $\geq 2.0$ g/dl  
3. Transfusion of $\geq 2$ units of packed cells or whole blood  
4. Need for reoperation  
5. Fatal | | |
|                            | **RECORD4**<sup>16</sup> Clinically overt bleeding:  
1. In critical organ; for example, intracranial, retroperitoneal, intraocular or intraspinal  
2. Fall of haemoglobin $\geq 2.0$ g/dl (calculated from postoperative level)  
3. Transfusion of $\geq 2$ units of blood  
4. Need for operation  
5. Fatal | | |
|                            | **ADVANCE**<sup>17</sup> Acute overt clinical bleeding with one of the following:  
1. Critical site (intracranial, retroperitoneal, intraocular, intraspinal, periocardial)  
2. Fall in haemoglobin $\geq 2$ g/dl within 24 h  
3. Transfusion $\geq 2$ units of packed cells | Not allowed |

Continued
<table>
<thead>
<tr>
<th>Study name</th>
<th>Major bleeding complication definition</th>
<th>Antiplatelet agent</th>
</tr>
</thead>
</table>
| ADVANCE 2     | 4. Need for surgical intervention at operative site  
5. Intramuscular bleeding with compartment syndrome  
6. Fatal                                             | Not allowed                            |
| ADVANCE 3     | As in ADVANCE 1                                                                                         | Not allowed                            |
| Treatment of venous thromboembolism                   |                                                                                                         |                                        |
| THRIVE        | Clinically overt bleeding:  
1. In critical sites  
2. Fall of haemoglobin ≥2.0 g/dl  
3. Transfusion of ≥2 units of blood or packed cells  
4. Fatal                                             | Aspirin at lowest effective dose permitted |
| RE-COVER      | Clinically overt bleeding:  
1. In critical sites  
2. Fall of haemoglobin ≥20 g/l  
3. Transfusion of ≥2 units of blood or packed cells  
4. Fatal                                             | Aspirin ≤100 mg daily permitted         |
| RE-SONATE     | Not stated                                                                                               | Not stated                              |
| REMEDY        | Not stated                                                                                               | Not stated                              |
| EINSTEIN      | Clinically overt bleeding:  
1. In critical sites; for example, intracranial and retroperitoneal  
2. Fall of haemoglobin ≥20 g/l  
3. Transfusion of ≥2 units of blood or packed cells  
4. Fatal                                             | Aspirin ≤100 mg daily or clopidogrel 75 mg daily, or both, were permitted |
| Prevention of embolic events in atrial fibrillation    |                                                                                                         |                                        |
| SPORTIF III   | Occurrence of at least one of the following:  
1. Intracranial, retroperitoneal, intraocular, intraspinal, pericardial or atraumatic intra-articular bleeding  
2. Clinically overt bleeding with fall of haemoglobin ≥20 g/l  
3. Transfusion of ≥2 units of erythrocytes or whole blood  
4. Fatal                                             | Aspirin ≤100 mg daily permitted (21%)*   |
| SPORTIF V     | As in SPORTIF III                                                                                       | As in SPORTIF III (18%)*                |
| RE-LY         | Occurrence of at least one of the following:  
1. Critical area or organ bleeding; for example, intracranial  
2. Clinically overt bleeding with fall of haemoglobin ≥20 g/l  
3. Transfusion of ≥2 units of blood  
4. Need for surgery  
5. Fatal                                             | Aspirin <100 mg daily or antiplatelet agent permitted (40%)* |
| ROCKET AF     | Clinically overt bleeding:  
1. In critical anatomic site; for example, intracranial, retroperitoneal, ocular, spinal, pericardial, articular or intramuscular with compartment syndrome  
2. Fall of haemoglobin ≥2.0 g/dl  
3. Transfusion of ≥2 units of whole blood or packed cells  
4. Permanent disability  
5. Fatal                                             | Aspirin ≤100 mg daily or monothienopyridine therapy permitted (38.5%)* |
| AVERROES      | Clinically overt bleeding:  
1. In critical sites; for example, intracranial, retroperitoneal, ocular, spinal, pericardial, articular or intramuscular with compartment syndrome  
2. Fall of haemoglobin ≥2.0 g/dl  
3. Transfusion of ≥2 units of packed cells  
4. Fatal                                             | Thienopyridine therapy permitted if needed |
| ARISTOTLE     | Clinically overt bleeding:  
1. In critical sites  
2. Fall of haemoglobin ≥2.0 g/dl over a 24 h period  
3. Transfusion of ≥2 units of packed cells  
4. Fatal                                             | Aspirin ≤165 mg daily or monothienopyridine permitted (32%)* |
patients with atrial fibrillation and not treated with aspirin.41 But the preliminary results from a substudy of the RE-LY trial did not show this relationship.42 Nonetheless, taken together, the differences in thrombotic, inflammatory and platelet pathways could have accounted for some of the differences in coronary events. Furthermore, there was discordance in the main findings of SPORTIF III25 and SPORTIF V26. Although both studies were similar in design, there were important dissimilarities. SPORTIF III25 was conducted in Europe, Asia plus Australasia and SPORTIF V26 was performed in North America. The design of the latter study26 was double-blind but SPORTIF III was an open-label trial.25 Of note, the primary endpoint, consisting of stroke and systemic embolism, was 2.3% per year for the warfarin group and 1.6% per year for the ximelagatan group in SPORTIF III.25 Conversely, it was 1.2% per year for the warfarin group and 1.6% per year for the ximelagatan group in SPORTIF V.26 There were also differences in the occurrence of major bleeding complications (figure 3A). The authors attributed the differences to better dose regulation, control of hypertension or hyperlipidaemia, other differences in patient characteristics or management or chance.26

Evaluation for a summarised risk for major bleeding complications among these studies has been challenging because of the marked variation in study protocol and endpoint definition (table 2). Although there was little difference in major bleeding complications for the four agents when compared with control, the rates were higher for rivaroxaban32 and apixaban33 34 in ACS patients, and influenced this outcome. Likely, several of these patients were receiving antiplatelet therapy, and probably treated with these two agents.

Indeed, major bleeding complication rates have been noted to increase by 40–70% among those receiving aspirin plus clopidogrel in the RE-LY trial.43 Majority of these ACS patients were receiving dual antiplatelet agents. Not surprisingly, when these trials were excluded from analysis, evidence for heterogeneity was lost. Therefore, extreme caution has to be exercised when considering combining antiplatelet and antithrombotic agents because of the high bleeding risk.

Despite differences in the risks for MI/ACS and major bleeding complications, all-cause mortality was lower among those treated with dabigatran, rivaroxaban and apixaban compared with control. Better survival was also observed among patients with ACS treated with oral anticoagulation. All-cause and vascular mortalities were significantly lower among those receiving moderate intensity of warfarin plus aspirin compared to aspirin alone.44 Part of the reason for lower mortality for patients treated with novel antithrombotic agents may be related to the lower rates of haemorrhagic stroke for those with atrial fibrillation.27 28 30 If this finding is real then it may supersede the shortfalls of these agents such as lack of antidote for reversal of effects and assay to determine its therapeutic efficacy.

There are several limitations in the study. Importantly, there were differences in study population, protocol and procedures. Duration of follow-up varied considerably across trials. Individual patient outcome information was also not available. Definitions of outcome measures varied considerably in the studies and there were subjective elements in adjudication, especially for bleeding complications. Furthermore, not all the outcomes were reported in every trial. Silent MI

### Table 2  Continued

<table>
<thead>
<tr>
<th>Study name</th>
<th>Major bleeding complication definition</th>
<th>Antiplatelet agent</th>
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<tbody>
<tr>
<td>Treatment of acute coronary syndrome</td>
<td>Occurrence of one of the following: 1. Bleeding in critical sites; for example, intracranial, retroperitoneal, ocular, spinal, pericardial, articular or intramuscular with compartment syndrome 2. Fall of haemoglobin ≥2.0 g/dl 3. Transfusion of ≥2 units of packed cells or whole blood 4. Fatal</td>
<td>All patients receiving dual antiplatelet agents</td>
</tr>
<tr>
<td>ATLAS ACS 2</td>
<td>Occurrence of one of the following: 1. Fall of haemoglobin ≥5.0 g/dl or haematocrit &gt;15% 2. Intracranial haemorrhage</td>
<td>All patients received low-dose aspirin and thienopyridine permitted</td>
</tr>
<tr>
<td>TIMI 512</td>
<td>Occurrence of one of the following: 1. Bleeding in critical sites; for example, intracranial, retroperitoneal, ocular, spinal, pericardial, articular or intramuscular with compartment syndrome 2. Fall of haemoglobin ≥2.0 g/dl 3. Transfusion of ≥2 units of packed cells or whole blood 4. Fatal</td>
<td>All patients received aspirin ≤165 mg daily and thienopyridine therapy permitted</td>
</tr>
<tr>
<td>APPRAISE</td>
<td>Occurrence of one of the following: 1. Fall of haemoglobin ≥5.0 g/dl or haematocrit &gt;15% 2. Intracranial haemorrhage</td>
<td>Use of aspirin and thienopyridine permitted</td>
</tr>
<tr>
<td>APPRAISE 2</td>
<td>Occurrence of one of the following: 1. Fall of haemoglobin ≥5.0 g/dl or haematocrit &gt;15% 2. Intracranial haemorrhage</td>
<td>Use of aspirin and thienopyridine permitted</td>
</tr>
</tbody>
</table>

*Proportion receiving antiplatelet therapy. Please refer to footnote of table 1 for acronyms.
may be actively sought out for in some studies, especially after revascularisation procedures, with routine electrocardiography or cardiac enzyme assays. But this approach may not be adopted in other trials. While this difference could have accounted for variation observed among studies, it was less likely to impact on the results within a study. Another limitation was that there was only one author in the study; there may be potential bias in study appraisal and selection stages. However, this concern is mitigated somewhat by relatively small total number of trials and fairly well-defined outcome measures. Nonetheless, these findings were instructive in providing insight on the relative occurrence adverse cardiovascular events impacting on the choice of these agents in specific patient subsets requiring anticoagulation. As with any results from meta-analyses, a firm conclusion can only be drawn from well-conducted, adequately powered randomised trials.

Competing interests None.

Provenance and peer review Not commissioned; externally peer reviewed.

Data sharing statement There are no additional data available.

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Oral novel antithrombotic agents and coronary risk

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Koon-Hou Mak

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