Economic evaluation of direct oral anticoagulants (DOACs) versus vitamin K antagonists (VKAs) for stroke prevention in patients with atrial fibrillation: a systematic review and meta-analysis

Rini Noviyani 1,2, Sitaporn Youngkong 1,3, Surakit Nathisuwan 4, Bhavani Shankara Bagepally 5, Usa Chaikledkaew 1, Nathorn Chaiyakunapruk 6, Gareth McKay 7, Piyamitr Sritara 8, John Attia 9, Ammarin Thakkinstian 1,10

Abstract

Objectives To assess cost-effectiveness of direct oral anticoagulants (DOACs) compared with vitamin K antagonists (VKAs) for stroke prevention in atrial fibrillation (AF) by pooling incremental net benefits (INBs).

Design Systematic review and meta-analysis.

Setting We searched PubMed, Scopus and Centre for Evaluation of Value and Risks in Health Registry from inception to December 2019.

Participants Patients with AF.

Main outcome measures The INB was defined as a difference of incremental effectiveness multiplied by willingness to pay threshold minus the incremental cost; a positive INB indicated favour treatment. These INBs were pooled (stratified by level of country income, perspective, time-horizon, model types) with a random-effects model if heterogeneity existed, otherwise a fixed effects model was applied. Heterogeneity was assessed using Q test and I² statistic. Risk of bias was assessed using the economic evaluations bias (ECOBIAS) checklist.

Results A total of 100 eligible economic evaluation studies (224 comparisons) were included. For high-income countries (HICs) from a third-party payer (TPP) perspective, the pooled INBs for DOAC versus VKA pairs were significantly cost-effective with INBs (95% CI) of $6632 ($2961.67 to $10 301.72; I²=59.9%), $6353.24 ($4076.03 to $8630.45; I²=59.9%), $7664.58 ($2979.79 to $12 349.37; I²=59.9%) and $8573.07 ($1877.05 to $15 269.09; I²=59.9%) for dabigatran, apixaban, rivaroxaban and edoxaban relative to VKA, respectively but only dabigatran was significantly cost-effective from societal perspective (SP) with an INB of $11 746.96 ($2429.34 to $21 064.59; I²=52.4%). The pooled INBs of all comparisons for upper-middle income countries (UMICs) were not significantly cost-effective. The ECOBIAS checklist indicated that risk of bias was mostly low for most items with the exception of five items which should be less influenced on pooling INBs.

Conclusions Our meta-analysis provides comprehensive economic evidence that allows policy makers to generalise cost-effectiveness data to their local context. All DOACs may be cost-effective compared with VKA in HICs with TPP perspective. The pooling results produced moderate to high heterogeneity particularly in UMICs. Further studies are required to inform UMICs with SP.

Summary box

What is already known about this subject?

► A large number of economic evaluation studies on direct acting oral anticoagulants (DOACs) and vitamin K antagonists (VKAs) were conducted in various healthcare settings to guide health policy makers in relation to reimbursement of DOACs.

► The previous systematic reviews that compared DOACs with VKAs for stroke prevention in atrial fibrillation did not provide an overall quantitative synthesis.

What are the new findings?

► This is the first quantitative meta-analysis of 100 economic evaluations (that included 144 comparisons) of all four DOACs with VKAs applying pooled incremental net benefit.

► Our findings indicated that DOACs might be significantly more cost-effective than VKAs in high-income countries using a third-party payer perspective while no DOACs were more cost-effective in upper-middle income countries (UMICs), regardless of any perspective was used.

► We found that country socioeconomic status and the methodological approach used potentially influenced the cost-effectiveness of DOACs compared with VKAs.
Evidence synthesis

Summary box

How might it impact clinical practice in the foreseeable future?

- While clinical efficacy and safety of DOACs over VKAs are established, these agents, at their current pricing, are cost-effective only in high-income countries but not in UMICs due partly to the lower socioeconomic status and the small number of studies available.
- Policy makers and pharmaceutical companies should together consider potential pathways to increase access to these useful agents by considering the impact of socioeconomic status on the cost-effectiveness for UMICs and potentially low-income and middle-income countries.

PROSPERO registration number CRD 42019146610.

Introduction

Atrial fibrillation (AF), the most common cardiac arrhythmia,1 is an important global health issue2 with an incidence of 596.2 cases/100 000 population in the Global Burden of Disease Study.3 Recent projections based on various national databases suggest that the incidence has doubled or tripled in the past decade.4-6 Complications of AF, particularly stroke, lead to significant morbidity and mortality.2 3 Disability-adjusted life years (DALYs) lost due to AF have increased almost linearly during the past 20 years, with a current global estimate of 5·98 million DALYs lost in 2017 alone.2

Oral anticoagulants such as vitamin K antagonists (VKAs, eg, warfarin) and direct oral anticoagulants (DOACs) are the cornerstone of stroke prevention in AF.7 VKAs have several limitations including the need for frequent monitoring as a consequence of numerous drug interactions.7 DOACs (ie, dabigatran, rivaroxaban, apixaban and edoxaban) were developed to reduce these limitations. Oral anticoagulants (VKAs, eg, warfarin) and direct oral anticoagulants (DOACs) are the cornerstone of stroke prevention in AF.7 VKAs have several limitations including the need for frequent monitoring as a consequence of numerous drug interactions.7 DOACs (ie, dabigatran, rivaroxaban, apixaban and edoxaban) were developed to reduce these limitations. Data from controlled trials and real-world studies suggest that DOACs are non-inferior to VKAs and have some advantages8-10 which has led to their recommendation over VKAs in the AF guidelines of many developed countries.11-15

Multiple cost-effectiveness studies have compared DOACs with VKAs in various healthcare settings to inform health policy including five systematic reviews (SRs) of economic evaluations.11-15 However, none have provided an overall quantitative synthesis of their findings. Recently, SR and meta-analysis (SR-MA) of economic outcomes have been performed by converting incremental cost–effectiveness ratio (ICER) to incremental net benefit (INB), and then pooling across studies.16 The ICER, estimated by dividing incremental cost with incremental effectiveness, could be interpreted that the intervention is said to be cost-effective if it is lower than the willingness to pay (WTP) threshold. However, the ICER is controversial in some state, that is, negative ICER may be due to a lower cost but higher effectiveness, or higher cost along with lower effectiveness of the intervention. Therefore, Crespo et al16 had suggested pooling the INB across studies, defined as a difference of incremental effectiveness multiplied by WTP threshold minus the incremental cost, which could be directly interpreted, that is, a positive INB indicated favour the intervention. This quantitative synthesis requires stratification by economic factors (eg, level of country income, time horizon, perspective, economic models and so on) to minimise heterogeneity.17-18 This SR-MA summarises the cost-effectiveness of individual DOACs compared with VKAs for stroke prevention in patients with AF to inform policy decisions in countries with limited resources.

Methods

This SR-MA was reported in accordance with the Preferred Reporting Items for Systematic Reviews and Meta-Analysis 2020 statement and the review protocol was registered at PROSPERO.19

Data sources and search strategy

We performed a comprehensive search in PubMed, Scopus and Centre for Evaluation of Value and Risks in Health (CEVR) databases from inception to 7 December 2019, see online supplemental appendix 1. Studies were selected if they met the following criteria included patients with AF, primarily/secondarily aimed to compare VKAs (ie, warfarin or acenocoumarol or phenprocoumon or coumarin) with DOACs (ie, dabigatran, apixaban, rivaroxaban and edoxaban), and reported ICER, quality-adjusted life years (QALYs) or INB. Studies were excluded if they provided insufficient data for synthesis.

Data extraction

Two investigators (RN and BSB) independently extracted data. Disagreement was resolved in consultation with senior authors (SY and AT). Extracted data included study characteristics, study population, interventions, economic data (ie, perspective, WTP threshold or gross domestic product estimates from the World Bank according to the study year, time-horizon, currency, economic model) and findings. In addition, data for pooling were also extracted including mean cost, incremental cost, clinical effectiveness, incremental effectiveness and ICERs together with SE, or 95% CI. Incremental costs and effectiveness were also extracted from the cost-effective plane using Web-Plot-Digitizer software V.4.2.20 21

Risk of bias

We assessed risk of bias for included studies using the economic evaluations bias (ECOBIAS) checklist.22 The first part evaluated the overall bias which consisted of the following 11 items: narrow perspective, insignificant comparator, cost measurement omission, intermittent data collection, invalid valuation, ordinal ICER, double-counting, inappropriate discounting, limited sensitivity analysis, sponsor and reporting/dissemination. The second part specifically evaluated risk of bias of the model specifications in economic evaluations consisting of three subdomains, that is, structure of the model (four items), data (six items) and consistency (one item). Each item was graded as yes, no, partly, unclear or not applicable, where yes and no referred to high and low risk of bias, respectively.

Data analysis

The primary outcome of interest was INB. Economic data were harmonised by converting all currency data using purchasing power parity for the year 2019.23 In addition, different scenarios were applied to estimate INB and its variance based on the methods suggested by Crespo et al16 (as follows: INB = K × ΔE − ΔC , or INB = ΔE × (K − ICER) where K is the WTP threshold, ΔC the incremental cost, ΔE the incremental effectiveness, ICER the incremental cost and incremental effectiveness ratio), and our expanded methods are published previously.17-18 see online
supplemental appendix 2. A positive INB indicated favouring treatment (ie, intervention is cost-effective), whereas a negative INB indicated favouring comparator (ie, intervention is not cost-effective).\textsuperscript{16,14,25} Heterogeneity was assessed using the Cochrane-\textit{Q} test and \textit{I}\textsuperscript{2} statistic and considered present if \textit{I}\textsuperscript{2} ≥25% or if the \textit{p} value was <0.1. The INBs were pooled across studies, stratified by country income (classified by the World Bank),\textsuperscript{16} time-horizon, economic model and perspective, using a random-effects model (Der Simonian and Laird method) if heterogeneity was present, or an inverse-variance model if not.\textsuperscript{25}

Meta-regression, sensitivity or subgroup analyses were undertaken to explore sources of heterogeneity such as discount rate, WTP threshold, data source and funding source. Publication bias was assessed using Egger’s test and funnel plots where number of studies/comparisons was 10 or more. Where a funnel plot was asymmetrical, a contour-enhanced funnel plot was constructed to assess if the asymmetry was due to missing studies or heterogeneity. All analyses were performed using STATA V.16. A two-sided \textit{p}<0.05 was considered statistically significant except for heterogeneity tests, in which \textit{p}<0.10 was used.

**Results**

**Study selection and characteristics**

Of the 1585 studies identified, 100 met the inclusion criteria. List of 14 excluded studies along with reasons are provided in online supplemental appendix 3 eTable 3.1. Of those, 86, 13 and 1 study were conducted in high-income countries (HICs), upper-middle income countries (UMICs) and low/middle income country, respectively. Comparisons included dabigatran versus warfarin (N=49),\textsuperscript{27–75} apixaban versus warfarin (N=39),\textsuperscript{28–30,32–41,43,45,51,62,65–67,69,72,73,76–92} rivaroxaban versus warfarin (N=34)\textsuperscript{28–30,32–38,40,41,43,45,50,51,57,62,65–69,72,73,83,93–101} and edoxaban versus warfarin (N=16)\textsuperscript{28,30,38,40,43,45,51,62,67,72,73,102–106} (see figure 1).

Characteristics are summarised in table 1 and online supplemental appendix 3 eTable 3.2. Most studies used a third-party payer (TPP) perspective (N=83),\textsuperscript{27–29,35–37,41,44,46–53,56–58,60–63,65–73,76–78,81–89,91–95,97,98,100,102,103,105–106} followed by societal perspective (SP) (N=21)\textsuperscript{28,33,36,43,54,57,59,64,72,73,80,99,101,104,113,125,127,128} and patient perspective (N=4).\textsuperscript{54,57,66,74} Most studies used Markov models and a lifetime-horizon with discounting for both cost and outcomes. About 90% of studies stated no conflict of interest, and 56% were funded by pharmaceutical companies.

Clinical and utility parameters were mostly taken from published literature. Country-specific and GDP-based thresholds were used for WTP in 73\textsuperscript{27–31,34–41,44–46,49,51,52,54,57–67,70,72,73,76–80,82–85,88,89,93,94,97,99–101,103–105,107–110,112–115,117–121,123–125,127,129} and 23 studies, 33\textsuperscript{43,45,50,55,56,68,69,71,74,75,81,86,87,91,95,99,100,102,103,105–126} respectively. Eighty-four studies with 166 comparisons\textsuperscript{29–32,34–41,44–50,52,54,56,57,60–64,66,67,69–71,73–85,87–89,91–94,96,97,99,100,102,104–129} reported increased cost-effectiveness with DOACs compared with warfarin/derivatives, in contrast to the remainder (58 comparisons from 24 studies) which did not.

**Risk of bias assessment**

Across all 22 items from the ECOBIAS checklist, 17 items where more than 70% of studies were graded as low risk of bias, see online supplemental appendix 3 eTable 3.3. Therefore, risk of bias was mostly low for most items with the exception of five items including narrow perspective, double-counting, inappropriate discounting, reporting and dissemination and internal consistency. However, these biases should be less influenced on pooling INBs because they were occurred in both intervention and control groups. Typically, the worst areas are described in figure 1. **Figure 1** Study selection flow. DOACs, direct oral anticoagulants; VKAs, vitamin K antagonists; CEVR, Centre for Evaluation of Value and Risks in Health databases.

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**Identification of new studies via databases and registers**

- Records identified from: Published N=355 Scopus N=950 CEVR, N=300
- Records removed before screening: Duplicate records (N=355) Records marked as ineligible by automated tools (N=0) Records removed for other reasons (N=0)
- Records excluded: Not interested target population (N=519) Not interested outcomes (N=368) Not DOACs nor VKAs (N=152) Cost analysis studies (N=4) Narrative reviews (N=2) Systematic reviews of clinical studies (N=12) Not cost-utility analysis (N=2)
- Records excluded: Not DOACs nor VKAs (N=5) Not interested outcomes (N=2) Narrative reviews (N=3) Conference abstract (N=2) Duplicated article (N=1) Could not retrieve full-text article (N=1)

**Risk of bias assessment**

Across all 22 items from the ECOBIAS checklist, 17 items where more than 70% of studies were graded as low risk of bias, see online supplemental appendix 3 eTable 3.3. Therefore, risk of bias was mostly low for most items with the exception of five items including narrow perspective, double-counting, inappropriate discounting, reporting and dissemination and internal consistency. However, these biases should be less influenced on pooling INBs because they were occurred in both intervention and control groups. Typically, the worst areas are described in figure 1. **Figure 1** Study selection flow. DOACs, direct oral anticoagulants; VKAs, vitamin K antagonists; CEVR, Centre for Evaluation of Value and Risks in Health databases.
Evidence synthesis

Table 1 General characteristics of the studies included (created by the authors)

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*The total number of studies are more than 100 because individual studies applied multiple methods.

comparator, thus, should be cancelled out when calculation of the INB (a ratio of an incremental cost and QALYs).

**Pooling of INB**

**Dabigatran versus VKAs**

Based on 40 studies with 48 comparisons in HICs with lifetime-horizon, the pooled INBs were $6432.70 from a TPP (95% CI $2961.67 to $10 303.72; I²=59.9%) and $11 746.96 from an SP (95% CI $2429.34 to $21 064.59; I²=52.4%). The corresponding pooled INBs in UMICs (nine studies with 13 comparisons) were $49 000.59 from a TPP (95% CI −$25 326.64 to $124 127.82; I²=99.8%) and −$14 709.67 from an SP (95% CI −$22 648.61 to −$6770.74; I²=69.2%). Dabigatran was cost-effective compared with VKAs in HICs, but not in UMICs (see figure 2, and online supplemental appendix eFigure 4.1–4.4). According to meta-regression for HICs, only funding source and WTP could partially explain heterogeneity for TPP and SP whereas heterogeneity in UMICs could not be explained (see online supplemental appendix eTable 4.1). Subgroup analysis by WTP <$50 000 and funding source from pharmaceutical companies showed that dabigatran was cost-effective compared with VKAs (online supplemental appendix 4 eFigure 4.5–4.6). Publication bias was done in the studies in HICs with TPP indicating no evidence of asymmetry, see online supplemental appendix eFigure 4.7.

**Apixaban versus VKAs**

Based on 31 studies (33 comparisons) in HICs, the pooled INBs were $6353.24 from a TPP (95% CI $4076.03 to $8630.45; I²=0%) and $1516.13 from an SP (95% CI −$22 648.61 to −$6770.74; I²=69.2%). Apixaban was cost-effective compared with VKAs in HICs with a TPP but not with an SP (see figure 2, and online supplemental appendix eFigures 5.1–5.4). According to meta-regression for UMICs, only discount rates for cost/utility and clinical data source could explain heterogeneity for a TPP whereas the other factors could not explain heterogeneity (see online supplemental appendix eTable 5.1, eFigure 5.5–5.8). There was no evidence of asymmetry using funnel plots and Egger’s tests for those studies in HICs with TPP, see online supplemental appendix eFigure 5.9.
Rivaroxaban versus VKAs

Based on 26 studies with 28 comparisons in HICs, the pooled INBs were $7664.58 from a TPP (95% CI $2979.79 to $12 349.37; I²=0%) and $10 345.74 from an SP (95% CI $14 615.67 to $36 153.02; I²=30.7%). The corresponding pooled INBs in UMICs (seven studies with 10 comparisons) were −$27 567.34 from a TPP (95% CI −$21 631.83 to −$7068.64; I²=68.3%) and −$14 350.24 from an SP (95% CI −$21 631.83 to −$7068.64; I²=68.3%). Rivaroxaban was cost-effective compared with VKAs in HICs with lifetime-horizon from TPP, but not from SP, see figure 2, and online supplemental appendix eFigure 6.1–6.4. Furthermore, rivaroxaban was significantly not cost-effective compared with VKAs in UMICs. According to meta-regression for UMICs with TPP, none of economic factors could explain heterogeneity (see online supplemental appendix eTable 6.1). There was no evidence of asymmetry for pooling INBs in HICs and TPP, see online supplemental appendix eFigure 6.5.

Edoxaban versus VKAs

Based on 13 studies with 15 comparisons in HICs, the pooled INBs (95% CI) were $8573.07 from a TPP (95% CI $1877.05 to $15 269.09; I²=0%). The pooled INBs in UMICs (three studies with five comparisons) were −$11 062.53 from a TPP (95% CI −$941 291.97 to $919 166.9; I²=0%) and −$15 547.36 from an SP (95% CI −$23 316.39 to −$7778.33; I²=51.3%). Edoxaban was cost-effective compared with VKAs from TPP only in HICs, but not cost-effective in UMICs in both TPP and SP, see figure 2, and online supplemental appendix eFigure 7.1–7.3. Source of heterogeneity could be not explored for pooling INBs in HICs and TPP, see online supplementary appendix eTable 6.1). There was no evidence of asymmetry for pooling INBs in HICs and TPP.
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Author affiliations
1Mahidol University Health Technology Assessment (MUHTA) Graduate Program, Mahidol University, Bangkok, Thailand
2Department of Pharmacy, Faculty of Mathematics and Natural Sciences, Udayana University, Bali, Indonesia
3Social and Administrative Pharmacy Division, Department of Pharmacy, Faculty of Pharmacy, Mahidol University, Bangkok, Thailand
4Clinical Pharmacy Division, Department of Pharmacy, Faculty of Pharmacy, Mahidol University, Bangkok, Thailand
5ICMR-National Institute of Epidemiology, Chennai, India
6Department of Pharmacotherapy, College of Pharmacy, University of Utah, Salt Lake City, Utah, USA
7Centre for Public Health, School of Medicine, Dentistry and Biomedical Sciences, Queen’s University, Belfast, UK
8Division of Cardiology, Department of Medicine, Faculty of Medicine, Ramathibodi Hospital, Mahidol University, Bangkok, Thailand
9School of Medicine and Public Health, Faculty of Health and Medicine, University of Newcastle, New South Wales, New South Wales, Australia
10Department of Clinical Epidemiology and Biostatistics, Faculty of Medicine, Ramathibodi Hospital, Mahidol University, Bangkok, Thailand

Contributors RN, BSB, SY and AT conceived and designed the work. RN and BSB did the analysis. RN wrote the first draft of the manuscript with input from SY and AT. All authors interpreted the data, provided critical revision for important intellectual content and approved the final version to be published.

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ORCID iDs
Rini Noviyani http://orcid.org/0000-0002-9306-2053
Sitaporn Youngkong http://orcid.org/0000-0002-2448-3954
Surakit Nathisuwon http://orcid.org/0000-0003-4828-4412
Bhavani Shankara Bagepally http://orcid.org/0000-0003-0856-767X
Usa Chaiakedkaew http://orcid.org/0000-0001-9457-9823
Nathorn Chaiyakunapruk http://orcid.org/0000-0003-4572-8794
Garecky kart http://orcid.org/0000-0001-8197-6280
Pyamitrat Sritara http://orcid.org/0000-0003-0509-8943
John Attia http://orcid.org/0000-0001-9800-1308
Ammarin Thakkistant http://orcid.org/0000-0001-9991-386X

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