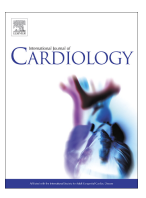
A review of global health technology assessments of non-VKA oral anticoagulants in non-valvular atrial fibrillation



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A Review of Global Health Technology Assessments of Non-VKA Oral Anticoagulants in

Non-Valvular Atrial Fibrillation

Running title: Review of Global HTA of NOACs in NVAF

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Abstract

Background: This review assessed global health technology assessment (HTA) reports and recommendations of non-vitamin K oral anticoagulants (NOACs) in non-valvular atrial fibrillation (NVAF).

Methods: HTA agency websites were searched for HTA reports evaluating NOACs versus NOACs or vitamin K antagonists. HTA methods and information on patient involvement/access were collected and empirically analyzed.

Results: The review identified 38 unique HTA reports published between 2012–2017 in 16 countries including 11 in Europe. NOACs that were cost-effective per local willingness-to-pay (WTP) thresholds were positively recommended for the treatment of NVAF. WTP thresholds ranged from €20,000 to 69,000. Apixaban was recommended in 10/12 (83%) countries, dabigatran in 9/13 (69%) countries, and rivaroxaban in 10/13 (76%) over warfarin. Edoxaban was recommended in 5/7 (71%) countries. Economic evaluations and recommendations comparing NOACs were sparse (two or three countries per NOAC) and generally favored apixaban and edoxaban, followed by dabigatran. Eleven HTA reports from four countries considered the patient voice (Canada [n=3], Scotland [n=3], England [n=4], Brazil [n=1]); however, only 2/11 (18%) developed recommendations based on this. Among the reports with a positive recommendation, 26/30 (87%) featured a decision that aligned with the approved regulatory label.

Conclusions: Most agencies recommended NOACs over warfarin for patients with NVAF. Few countries made statements recommending one NOAC over another. Given different WTP thresholds, a drug that is cost-effective in one market may not be in another. Therefore, the

various NOAC recommendations from HTA agencies cannot be generalized across different countries.

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1 Introduction¹⁰

1.1 HTA assessments across Countries

Health technology assessment (HTA) is a systematic evaluation of treatments to inform health policy, access and reimbursement decision-making.[1]. There are more than 40 countries with HTA agencies generating HTA recommendation reports in their respective markets [2,3]. Different HTA archetypes exist across countries; these are defined by the focus of their assessment (i.e., clinical and/or economic evidence), methods employed, submission processes and requirements, payment/reimbursement systems, national or regional assessments, and other pricing and pharmacoeconomic factors [4]. Prior research found that differences in assessment methodologies, mandates and political systems across countries can lead to variations in final recommendations for new drugs [5,6]. To this end, there has been a recent emphasis on a need for more standardized practices in HTA [6,7].

¹⁰ Abbreviations: AF, atrial fibrillation; BIM, budget impact model; CADTH, Canadian Agency for Drugs and Technologies in Health; CEA, cost-effectiveness analysis; CMA, cost-minimization analysis; CUA, cost-utility analysis; HAQI, Healthcare Access and Quality Index; HTA, health technology assessment; HTAi, Health Technology Assessment International; INAHTA, International Network of Agencies for Health Technology Assessment; ISPOR, International Society for Pharmacoeconomics and Outcomes Research; LFN, Pharmaceutical Benefits Board (Sweden); NICE, National Institute for Health and Care Excellence (United Kingdom); NMA, network meta-analysis; NOAC, non-vitamin k antagonist oral anticoagulant; NVAF, non-valvular atrial fibrillation; PBAC, Pharmaceutical Benefits Advisory Committee (Australia); RCT, randomized controlled trial; RWE, realworld evidence; SE, systemic embolism; VKA, vitamin k antagonist; WTP, willingness-to-pay.

1.2 NOACs for Treatment in NVAF

Global assessment is needed for treatments that are approved worldwide to shed insight on potential variation across countries in evaluation and approval of the same treatments. This is particularly pertinent for non-vitamin K antagonist oral anticoagulants (NOACs), specifically apixaban, dabigatran, edoxaban, and rivaroxaban, as they are increasingly being approved and recommended worldwide as treatment for stroke prevention in patients with NVAF [8]. Randomized controlled trials (RCTs) demonstrated that NOACs have similar or better efficacy and safety in comparison to vitamin k antagonists (VKAs) [9-12]. Differently from warfarin, NOACs can be administered without routine monitoring of anticoagulant levels. Despite the clinical advantages of NOACs, warfarin is still widely used in clinical practice, likely due to its established familiarity and lower cost [13].

Although clinical efficacy and cost-effectiveness of NOACs have been assessed [14], no comprehensive review of HTA reports assessing different NOACs exists. Most NOACs have been in the market for several years, while edoxaban was authorized more recently. Some differences in HTA submission methods may have occurred over this broad time horizon, resulting in different recommendations. Assessing the timing of NOAC submissions may reveal whether factors such as patient voices, real-world evidence (RWE), and data from more mature RCTs had an impact in later submissions compared to earlier ones. To advance this research, we performed a review of global HTAs that evaluated NOACs for treatment of patients with NVAF. We aimed to evaluate similarities and differences across country-level HTAs in methodology, data considerations, final decisions on recommended use of each NOAC, and preferential

statements for the NOACs (e.g., related to subgroups, dose, and clinical outcomes). This review addresses two research questions:

- What was the global clinical and economic value of the NOACs across national HTAs, and what methods, perspectives, and evidence were considered in the evaluation of such value?
- 2. How closely did HTA decisions and recommendations match the approved regulatory label and results of economic evaluations? If the decisions did not match the approved regulatory label, what factors led to the decisions?

2 Methods

2.1 Data Sources, Searches, and Identification of Studies

The search methods used to identify eligible HTA reports consisted of two phases. In phase 1, the websites of INAHTA, HTAi vortal, ISPOR, and the European Network for Health Technology Assessment were searched in August–September 2018 to identify countries and agencies that produce HTA reports (Supplemental Table 1). In phase 2, websites of each HTA agency identified were searched for publicly available HTA documents related to NOACs for prevention of stroke in patients with NVAF (Table 1), using keywords "atrial fibrillation," "oral anticoagulants," "apixaban," "dabigatran," "edoxaban," "rivaroxaban." No geographic, language, or temporal limits were applied to the search.

Documents eligible for inclusion were HTA reports of NOACs for treatment of patients with NVAF. Two investigators independently reviewed the identified documents to determine their eligibility for inclusion in the review. Any discrepancies were resolved by a third independent

investigator. Published HTA reports for a given country that evaluated only dabigatran or rivaroxaban were excluded unless such reports evaluating apixaban and edoxaban were also available for the same country; this ensured that the comparison of the four NOACs was similar across countries that evaluated multiple NOACs.

2.2 Data Extraction and Synthesis

One investigator independently extracted key information from the reports and a second investigator validated data for accuracy. Discrepancies between investigators were resolved by a third, independent investigator. If multiple reports were identified for a single HTA submission, they were extracted as a single report. In cases where updates to HTA reports were available, the more recent applicable evidence took precedence over older documents when summarizing main conclusions. However, all documents were considered in the evaluation of the methodology, results, and conclusions of the HTA reports. Extracted data elements are summarized in Supplemental Table 2. Due to the absence of a standard quality assessment instrument for HTA reports, no formal quality assessment was undertaken in this review.

A qualitative synthesis was performed to summarize key findings and patterns across HTA reports and identify gaps. The synthesis was conducted following an *a priori* framework with information clustered by country, HTA agency, type of NOAC, and type of evidence (clinical and/or economic) presented in the reports (Supplemental Figure 1). Addressing the first research question involved synthesis of clinical evidence, including the methods, results and conclusions of RCTs, network meta-analyses (NMAs), and RWE. Additionally, economic evidence, including the methods, results and conclusions of economic evaluations, were synthesized. The cost-effectiveness results and willingness-to-pay (WTP) thresholds were converted to 2019 Euros for comparability. The cost conversions were completed by first inflating values to the

year 2019 and then using country-specific conversion rates to convert each currency to Euro [15-20]. In answering the second research question, the final recommendations of the assessments, drivers of decisions, and approved regulatory labels reported in each HTA report were collated and compared. In addition, comments from patients and patient representatives considered in HTAs were categorized and compared.

3 Results

3.1 Search Results

Phase 1 searches yielded websites for 68 agencies across 36 countries (Supplemental Table 1). The phase 2 search of agency websites yielded 8,886 records. Results by country appear in Supplemental Table 3. Fifty HTA reports (38 unique and 12 related documents) from 16 countries were included in this review (Supplemental Figure 2). Despite no date limit set on the search strategy, the publication date of included reports ranged from 2012 to 2017. This aligns with the approval date of NOACs (2011-2015). Among the reports, four were from Netherlands, four Sweden, two Colombia, two Poland, one Spain and one Brazil.

3.2 HTA Report Characteristics

Among the 16 countries with HTA reports providing clinical and/or economic results for NOACs, all (100%) reported on apixaban, 15 (94%) on dabigatran and rivaroxaban, and 10 (63%) on edoxaban. Publication years ranged from 2011 to 2018, with 52% published in 2012 and 2013. All HTA reports included evidence on stroke and major bleeding, and all but one reported evidence for systemic embolism (SE) [21,22]. Eleven of 16 (69%) countries were European; the remaining countries were Canada, Brazil, Colombia, Australia, and Singapore. Thirteen (81%) countries provided both clinical and cost-effectiveness results (in one or across

multiple reports); HTA documents for Germany, France, and Spain reported only clinical results. Reports from only four countries (25%) considered patient voice [23-33]. The characteristics of included HTA reports by country are summarized in Table 1.

3.3 Clinical and Economic Value (Research Question 1)

3.3.1 Clinical Evidence

All 38 HTAs reported clinical evidence, from either RCTs (n=38), NMAs of RCTs (n=26), or RWE (n=8; Table 1). None of the RCTs compared two or more NOACs directly. Hence, HTA agencies compared two NOACs using RCTs comparing NOACs with vitamin K antagonists or aspirin in an NMA.

Among the 26 HTAs presenting NMA results, six (26%) reported one NOAC's clinical superiority over another [21,25,34-37]. For primary outcomes (stroke, SE, and/or major bleeding), apixaban had significantly better clinical efficacy and safety compared with dabigatran (n=4), rivaroxaban (n=2), and edoxaban (n=1), and dabigatran had significantly better efficacy and safety than rivaroxaban. Edoxaban had significantly lower major bleeding risk compared with dabigatran (n=1) and rivaroxaban (n=2). No HTA reports showed rivaroxaban to be clinically superior to another NOAC.

RWE was sparsely considered in HTA reports. Two reports included RWE in the clinical inputs, but there was no evidence suggesting the additional RWE impacted the main conclusions of the reports.[38,39]

3.3.2 Economic Evidence

Among the 35 reports examining NOACs' cost-effectiveness, the most commonly used methods were cost-utility analysis (CUA) (n=18) and cost-effectiveness analysis (CEA) (n=19), alone or

in conjunction with other methods (Table 2). Among these, all reported QALY estimates and four of the CUAs (Canada, Norway, and Scotland) reported QALYs and cost per life-year [30,40-42]. Cost per life-year was also reported by two reports that used both CEA and CUA from the Netherlands [43,44]. Four reports presented information from cost-minimization analyses (CMAs) (Australia [n=2], Singapore, and Sweden). CEA and CMA were both presented in reports from Australia and Singapore [45-48]. Eleven HTAs (six countries) reported WTP thresholds; a drug that was not cost-effective based on these thresholds was not recommended by the country's agency. The WTP was comparable across countries and agencies. The thresholds ranged from 20,000 to 50,000 in United States dollars, British pounds, or Euros (Table 2). When converted to Euros, the range was still 20,000 to 50,000 with the exception of one report from Norway that gave a WTP threshold of 580,000 Norwegian krone (€68,963) [41]. Of note, the review included four reports from the Netherlands, two assessing clinical and economic value (dabigatran and rivaroxaban)[43,44] and two reporting clinical value only (apixaban and edoxaban) [36,49]. The clinical value only reports (Germany [50], France [35], and the Netherlands [apixaban and edoxaban]) were not considered for economic value or final recommendations.

3.3.3 Budget Impact Models

Thirty-five reports presented economic evidence, 13 of which included a budget impact model (BIM) (see Supplemental Table 3) [21,23,28-30,34,37,41,42,47,51-53]. Australia and Ireland were the only countries with reports comparing NOACs directly in the BIMs. The remaining countries compared NOACs with warfarin and/or aspirin.

3.3.4 HTA Recommendations

3.3.4.1 NOACs vs. Warfarin

Most NOACs were considered cost-effective compared with VKAs, based on findings from 14 countries. Of the countries that evaluated cost-effectiveness, most found rivaroxaban (10/13), apixaban (8/12), dabigatran (8/13), and edoxaban (5/7) to be cost-effective over VKAs (Table 3).

3.3.4.2 NOACs vs. other NOACs

Four of 12 countries providing recommendations for apixaban (Canada, Ireland, Norway, and Scotland) reported cost-effectiveness comparisons between apixaban and other NOACs. Rivaroxaban was least cost-effective, as only two countries showed rivaroxaban to be as costeffective as another NOAC (dabigatran) but was not as cost-effective as apixaban or edoxaban (Table 3). Apixaban, dabigatran and edoxaban were each as or more cost-effective than other NOACs in 2-3 countries (Table 3).

3.3.5 Assessment of HTA Methods and Approved Regulatory Label Alignment (Research Question 2)Patient Voice

Eleven HTA reports across four countries considered the patient voice as part of their assessment (Canada [n=3 (27%)], Scotland [n=3 (27%)], England [n=4 (36%)], Brazil [n=1 (9%)]) [23-33]. The comments from patients and patient representatives were qualitatively synthesized. Specific inconveniences associated with warfarin included frequency of INR monitoring appointments, which affect day-to-day life and result in lost work time, and the concerns of food-drug, food-alcohol, and drug-drug interactions, which limit social activities and quality of life. Five HTA reports noted that patients or patient representatives expected that NOAC(s) evaluated would improve quality of life and/or provide relief from the burden of warfarin. Two of the nine HTAs

specifically stated that patient data were considered in reaching the final decision about treatment.

3.3.6 Patient Access

3.3.7 Alignment with Approved Regulatory Label

Of the 38 HTA reports, 14 did not report information on approved regulatory labels. Scotland was the only country that reported the date of the approved regulatory label. To address this missing information, individual agency sources were searched. A summary of the alignments between HTA recommendations and approved regulatory labels appears in Table 4.

Twenty-six of 30 reports with a positive NOAC recommendation aligned with the approved regulatory label, while four did not. In these cases, the HTA report recommended the NOAC for higher-risk group individuals based on CHADS₂ score cutoff, but the labels did not reflect that limitation. In two Canadian HTA reports, the recommendation applied to patients with CHADS₂ score ≥ 1 , while the regulatory label specified patients with a CHADS₂ score ≥ 2 [40]. In one report from France (clinical only) [35], edoxaban was recommended as a secondary treatment in patients with contraindication, low tolerance, or inability to achieve INR targets with VKAs. In contrast, the approved regulatory label included a broader population of patients with NVAF. A report from Poland recommended dabigatran for a narrower NVAF patient population (CHADS₂ score ≥ 3 and patient ages ≥ 75) compared to the approved regulatory label was wider [54].

3.3.8 Changes to HTA Recommendations over Time

Recommendation reports from five countries (Australia, Colombia, Canada, England, and Sweden) were updated or included an addendum. The drivers of such changes were additional

sensitivity analyses and variable adjustments and/or additional evidence that were not available at the time of original publication. Three reports (Australia, Canada, England) included more recent published reports to be more inclusive of the NOACs based on market availability In England, the original report (2012) concluded that rivaroxaban was cost-effective versus warfarin [55], whereas the updated documents (2015–2017) with evaluations of all four NOACs, determined that evidence was insufficient to justify conclusions of superiority among the NOACs [25-27]. Similarly, for Canada, one HTA report (2013) did not include apixaban, but the updated report (2017) included data from all four NOACs [24]. The update recommended NOACs over warfarin in patients with a CHADS₂ score ≥ 1 and concluded there was insufficient evidence to decide superiority among the NOACs. In Australia, Colombia, and Sweden, the addenda were driven by requests to evaluate additional variables and perform supplementary analyses to the original report(s) but the additions did not alter the recommendations of each report.

3.3.9 Decision Drivers

Decision drivers were usually not explicitly stated in the HTA reports, rather most countries included reasons or rationales for the decisions (Table 4). Sixteen reports concluded that cost-effectiveness was the reason for positive recommendations: three each from Australia [45,46,51] and Canada [56,57]; two each from England [55,58], the Netherlands [43], Norway [41], Scotland [30,42], and Sweden [38,59]. NICE CEA results specifically stated the drivers, which included discontinuation rates on the first line of treatment (apixaban) [25], lower rates of myocardial infarction, intracranial hemorrhage, and other clinically relevant bleeding (apixaban), and hemorrhagic stroke (edoxaban) [26]. Reports from Singapore and Columbia (apixaban, dabigatran, and rivaroxaban only) reported the reason for the reject decisions was the high cost

of the NOACs [21,22,47]. Drivers or reasons for decisions were not reported or not publicly available in 13 reports [28,29,34,36,39,45,48,49,52-54,60,61].

4 Discussion

This global review on NOACs for patients with NVAF yielded 38 unique HTA reports (50 documents; 16 countries). Most HTA reports recommended NOACs; few countries recommended one NOAC over another. This can be attributed to the clinical evidence, which was based on indirect comparisons between NOACs (mainly via warfarin) due to the lack of RCTs directly comparing individual NOACs. Cost-effectiveness was a major driver of positive recommendations; if a drug was not cost-effective based on local WTP thresholds, it was not recommended by the respective HTA agency.

Generally, NOACs were recommended by most agencies/countries. Exceptions included Colombia and Brazil, where apixaban, dabigatran, and rivaroxaban were not recommended, mainly due to high drug costs.[22,23] Brazil's HTA agency was concerned with uncertainty around existing trial data and lack of patient monitoring (absence of INR monitoring). Similarly, Singapore did not recommend dabigatran, based on unacceptable cost-effectiveness and budget impact results.[47] In contrast, the high cost of NOACs did not negatively impact results in Canada, Scotland, or England, where all NOACs were recommended over warfarin.

Healthcare cost, quality, and affordability/access could represent potential drivers of HTA decisions across countries and determine drug value. A systematic analysis from the Global Burden of Disease Study in 2016 evaluated healthcare access and quality globally by calculating a Healthcare Access and Quality Index (HAQI) [62]. The HAQI scores range from 0 to 100, with 100 representing the highest-quality healthcare. All countries in the review had high HAQI

scores (>90) except for Poland, Colombia, and Brazil. Colombia and Brazil are two of the countries in our review that did not recommend NOACs (HAQI scores of 62 and 64, respectively).

Canada, Scotland, and England were the only countries whose reports considered the patient voice; however, only one report from England incorporated such data into the final decision [23-33]. The few reports in this review that considered patient voice concluded there was an inconvenience and burden associated with warfarin that was not considered in the RCTs evaluating clinical efficacy and safety.

RWE was also sparsely considered in the HTA reports or updates. Three countries (Sweden, Australia, and Spain) mentioned the RELY-ABLE study, an observational follow-up of the RE-LY trial [63]. Sweden was the only country with reports including additional RWE in the clinical evidence. A potential reason for the lack of RWE in the NOAC HTAs is the year of publication. A limited number of HTA reports (n=11) were published in 2016 or later. Recent systematic literature reviews of RWE in NOACs have identified an absence of comparative RWE among the NOACs, particularly prior to 2015 [64,65]. Additionally, the use of RWE in HTAs is inconsistent highlighting a need for a policy on RWE [66]. Only in the last few years has the inclusion of RWE in HTAs gained traction.

Each HTA agency has a different drug-implementation program, with varying systems, regulations, and drug-approval processes related to patient access. One of the objectives of this review was to gain insights into global variations in patient access and implementation of HTA recommendations. However, information on these topics was rarely included in the HTA reports.

When evaluating discordance between HTA agency recommendations, it is useful to understand commonalities and differences between HTA agencies. For example, HTA agencies for Canada, Scotland, and England have much in common regarding how they evaluate new treatments and value the opportunity for early engagement with companies targeting their markets—NICE and CADTH recently launched a new collaboration to offer parallel scientific advice to the life sciences industry [67]. Hence, it is not surprising that these agencies would align in their NOAC recommendations. In contrast, as highlighted by a case study of HTA systems in Australia, Canada, England and Scotland, these four countries provided divergent recommendations using similar rationale and information.[6]. The variation in consideration of varying factors, during the decision-making and recommendation process by the agency could be a driver of conflicting conclusions based on similar evidence [6]. In the present review, similar evidence for clinical efficacy and safety of the NOACs was used across the HTA reports, but a few countries did not recommend the use of the NOACs. The differences in scientific standards, country-specific considerations, and variation in agency consideration could be a factor in some of the differences seen in the recommendations across agencies.

Finally, this review has several limitations. The search covered a wide range of global HTA bodies with no language limits. For search and screening, when available, native speakers were utilized to search for non-English reports on agency websites, as the websites need to be searched manually and cannot be searched with search strings like electronic databases. However, if a native language speaker was not available, searchers and reviewers relied on translation tools, which are not always accurate and may have caused some reports to be missed. However, if any potentially relevant reports were identified by automated translation tools during screening, they were included for full translation. Another limitation was that information in

HTA reports did not comprehensively address the research questions related to patient access. For example, many reports did not state dates of approved regulatory labels, so dates were collected from agency websites. Judgment calls were made on whether the recommendations matched the approved regulatory label. An additional limitation of this review is that data collection was based on public availability of information, which varies by country and agency. Countries like England, Australia, and Canada have a plethora of documents supporting HTA reports online with final decisions. In contrast, some countries, such as Poland, concealed methodological details and results from the publicly available version, limiting our access to comprehensive data.

5 Conclusions

The present review furthered the existing research in assessing HTA methods and variation in HTAs across countries worldwide. Through the evaluation of HTA reports on NOACs for the treatment of NVAF, we observed differences in methods and processes, such as methodology, patient involvement and included NOACs. However, only a portion of the differences across HTAs can be evaluated based on the report information. Other factors and data sources should be taken into account to gain a systemic understanding, such as agency regulation, healthcare systems, socioeconomic status, and political climate. Given the variation in HTA methodology across countries and the multifactorial influence on drug recommendations, differences in recommendations across various HTA agencies should be assessed considering the above factors and should not be generalized across different countries.

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Table 1. HTA Characteristics by Country (n=16)

Country,									. .	
Year* (# of reports)	Agency	Α	D	E	R	VKA	ASA	Clinical Evidence	Economic Evidence	Patient Input
Australia, 2013 (7)	PBAC	~	~		~	~	~	RCT, NMA of RCTs, RWE	CEA, CUA, CMA	No
Belgium, 2017 (1)	KCE	~	~	~	~	✓		NMA of RCTs	SLR of CEAs	No
Brazil, 2016 (1)	CONITEC	~	~		~	~		RCT, NMA of RCTs	Cost comparison	Yes
Canada, 2013** (5)	CADTH	~	~	~	~	~	~	RCT, NMA of RCTs	CUA	Yes
Colombia, 2016 (2)	IETS	~	~		~	~		RCT, NMA of RCTs	CEA	No
England, 2017 (5)	NICE	~	~	~	~	~	~	RCT, NMA of RCTs, RWE	CEA, CUA	Yes
France, 2016 (1)	HAS	~	~	~	~	√	2	RCT, NMA of RCTs	NA	No
Germany, 2013 (1)	IQWiG	~				~	~	RCT	NA	No
Ireland, 2013 (3)	NCPE	~	~		\checkmark	~	~	RCT, NMA of RCTs	CEA	No
Netherlands, 2015 (4)	GVS, CVZ, CFH	~	~	~	~	~		RCT	CUA	No
Norway, 2013 (1)	NoMA	~	~		~	~		RCT, NMA of RCTs, other HTAs	CUA	No
Poland, 2013 (2)	AOTMiT	~	~	~	~	~	~	RCT, NMA of RCTs	CUA	No
Scotland, 2015 (4)	SMC	~	~	~	~	~	~	RCT, NMA of RCTs	CUA	Yes
Singapore, 2018 (1)	ACE	~	~		~	~		RCT	CEA, CMA	No
Spain, 2016 (1)	AEMPS	~	~	~	~	~		RCT	NA	No
Sweden, 2016 (4)	TLV	V CE -	~	~	~	✓		RCT, NMA of RCTs, RWE	CEA, CMA	No

Abbreviations: A = apixaban; ACE = Agency for Care Effectiveness; AEMPS = Agencia Española de Medicamentos y Productos Sanitarios; AOTMiT = Agency for the Assessment of Medical Technology and Tariffs; ASA = aspirin; CADTH = Canadian Agency for Drugs and Technologies in Health; CEA = cost-effectiveness analysis; CFH = Committee of Pharmaceutical Aid; CMA = cost-minimization analysis; CONITEC = National Committee for Technology Incorporation; CUA = cost-utility analysis; CVZ = Health Care Insurance Board; D =

dabigatran; E = edoxaban; GVS = Medicine Reimbursement System; HAS = Haute Autorité de Santé; IETS = Institute of Technological Evaluation in Health; IQWiG = Institute for Quality and Efficiency in Health Care; KCE = Belgian Health Care Knowledge Centre; NA = not applicable; NCPE = National Centre for Pharmacoeconomics; NICE = National Institute for Health and Care Excellence; NMA = network meta-analysis; NoMA = Norwegian Medicines Agency; PABC = Pharmaceutical Benefits Advisory Committee; R = rivaroxaban; RCT = randomized controlled trial; RWE = real-world evidence; SLR = systematic literature review; SMC = Scottish Medicines Consortium; TLV = Dental and Pharmaceutical Benefits Agency; VKA = vitamin K antagonist *When more than one report was published for a given country, the characteristics were combined, and the most recent date was reported.

**All reports in 2013, except edoxaban report in 2017.

			,		- I		
Country , Year	Analysi s Time horizon	WTP	Compari son	Summary Metric	Summary Metric in 2019 Euros	Preferenti al Statement	
	CE A		A, D, R vs. warfarin	A\$45,000– 75,000/QALY	€32,395- 53,991/QAL Y ¹	Committee agreed to substitution of	
, 2013 10-20		and CMA NR 10–20		NR	NA	warfarin, aspirin, and potentially from no treatment with NOACs.	
Australia , 2011	CEA Lifetim e	NR	D vs. warfarin	A\$15,000- 45,000/QALY	€10,798- 32,395/QAL Y ¹	Dabigatran 150 and 110 mg recommen ded based on acceptable cost- effectivene ss	
Australia , 2013	CUA NR	NR	R vs. warfarin	A\$15,000– 45,000/QALY	€10,357- 31,072/QAL Y ¹	Rivaroxaba n recommen ded based on cost- effectivene ss compared with warfarin.	
Belgium, 2017	SLR of CEAs NR	NA	NOACs vs. VKAs	NA		NOACs were considered cost- effective against VKAs	
Brazil, 2016	Cost compari son	NA	A, D, R vs. warfarin	NR		The committee concluded	

Table 2. Characteristics	Methods	and Economic	Results of HTA	Reports
$1 \text{ abit } \underline{2}$. Characteristics	, muuus	, and Economic	NUSUIUS OI II I A	i incipul is

Country , Year	Analysi s Time horizon	WTP	Compari son	Summary Metric	Summary Metric in 2019 Euros	Preferenti al Statement
	NA					that a disadvanta ge of NOACs are its higher costs.
			A vs. warfarin D 150 vs.	C\$24,312/QALY C\$17,525/QALY	€17,542/QA LY ² €12,645/QA LY ²	The committee
Canada,	CUA 30–40	NR	warfarin D 110 vs. warfarin R vs.	C\$96,026/QALY	LY ² €69,287/QA LY ² €40,231/QA	concluded that the relative
2013	years (lifetim e)	INK	k vs. warfarin A vs. D	C\$55,757/QALY NR	LY ² NA	cost- effectivene ss of the
			A vs. R A, D, R vs. warfarin	NR NR	NA NA	- NOACs is uncertain.
Canada, 2017	CUA Lifetim e	NR	E vs. warfarin	C\$12,672/QALY	€8,702/QAL Y ²	Edoxaban was cost- effective compared with warfarin.
			E vs. R	NR		Dominated
		5	A vs. warfarin	COL\$97,501,541/ QALY	LY ^{3,4}	The committee
	CEA		D 150 vs. warfarin	COL\$74,462,000/ QALY	€22,605/QA LY ^{3,4}	concluded that costs of the
Colombi a, 2016	Lifetim e	NR	R vs. warfarin	COL\$91,981,682/ QALY	€27,924/QA LY ^{3,4}	NOACs were three
	C		D vs. R D vs. A	NR NR	NA NA	times Colombia's GDP per capita.
England,	CEA	NR	D 150 vs warfarin	£7645/QALY	€9,516/QAL Y ⁵	The NOACs
2015	30 years		A vs. warfarin	£9383/QALY	€ 11,679/QAL	strictly dominated

Country , Year	Analysi s Time horizon	WTP	Compari son	Summary Metric	Summary Metric in 2019 Euros	Preferenti al Statement
			E vs. warfarin	£12,881/QALY	$\begin{array}{c} Y^5\\ \hline \\ \hline \\ 16,033/QAL\\ Y^5 \end{array}$	over warfarin, but evidence is
			D 110 vs. warfarin	£13,565/QALY	€ 16,884/QAL Y ⁵	insufficient for cost- effectivene
			R vs. warfarin	£28,180/QALY	€ 35,075/QAL Y ⁵	ss among the NOACs.
England, 2017	CEA Lifetim e	£20,000/QA LY	NOACs vs. warfarin	NR	NA	Use of NOACs may be cost- effective compared with warfarin.
	CUA Lifetim e	£20,000; £30,000 (€26,710; €40,065)	D vs. warfarin	£18,900/QALY	€25,241/QA LY ⁵	Dabigatran was cost- effective compared with warfarin.
England, 2012			R vs. warfarin	£18,883/QALY	€25,218/QA LY ⁵	Rivaroxaba n was more
	CEA Lifetim	£20,000; £30,000	D vs. warfarin	£34,680	€46,315/QA LY ⁵	cost- effective
	e	(€26,710; €40,065)	R vs. D	NR	NA	than dabigatran and warfarin
France, 2016	NA	NA	NA	NA	NA	NA
Germany , 2013	NA	NA	NA	NA	NA	NA
T 1 1		€45,000/QA LY	A vs. warfarin	€23,669/QALY	€24,177/QA LY ⁵	Apixaban was cost-
Ireland, 2013	CEA NR	(€45,967/Q ALY)	A vs. D A vs. R	NR NR		effective compared with dabigatran,

Country , Year	Analysi s Time horizon	WTP	Compari son	Summary Metric	Summary Metric in 2019 Euros	Preferenti al Statement
						rivaroxaba n, and warfarin.
Ireland, 2012	CEA 30 years	€20,000– 30,000/QAL Y	R vs. warfarin	€22,663/QALY	€23,219/QA LY ⁵	Rivaroxaba n is not cost- effective compared with warfarin.
Ireland, 2011	CEA NR	€20,000– 30,000/QAL Y	D vs. warfarin	<80 years: €6,311/QALY 80 years or older: €20,654/QALY	<80 years: €6,492/QAL Y ⁵ 80 years or older: €21,246/QA LY ⁵	Dabigatran may be cost- effective compared with warfarin in patients with risk factors, but models contain uncertainti es.
		NR	D vs. warfarin	€7,719	€ 8,275	Dabigatran is cost- effective compared with warfarin.
Netherla nds, 2012	CUA Lifetim e	€20,000/QA LY	R vs. warfarin	€11,396/QALY	€12,217/QA LY ⁵	Rivaroxaba n is cost- effective compared with warfarin.
		NR	D vs. R	NR	NA	Dabigatran is interchange able with rivaroxaba n.

Country , Year	Analysi s Time horizon	WTP	Compari son	Summary Metric	Summary Metric in 2019 Euros	Preferenti al Statement
			R vs. warfarin	CHADS2-VASc = 1 and HAS- BLED = 0: NOK 317,550/QALY CHADS2-VASc = 2 and HAS- BLED = 1: NR	€37243/QA LY ⁶	
Norway, 2013	CUA Lifetim e	NOK 588,000/QA LY (€68,963)	D 150 vs. warfarin A vs. warfarin	CHADS2-VASc = 1 and HAS- BLED = 0: NOK 328,174/QALY CHADS2-VASc = 2 and HAS- BLED = 1: NOK 106,142/QALY CHADS2-VASc = 1 and HAS- BLED = 0: NOK 881,627/QALY CHADS2-VASc = 2 and HAS- BLED = 1: NR	CHADS2- VASc = 1 and HAS- BLED = 0: ϵ 38489/QA LY ⁶ CHADS2- VASc = 2 and HAS- BLED = 1: ϵ 12449/QA LY ⁶ CHADS2- VASc = 1 and HAS- BLED = 0: ϵ 103400/QA LY ⁶ CHADS2- VASc = 2 and HAS- DLED = 1	All NOACs were cost- effective compared with warfarin for patients of medium to high risk of stroke. Dabigatran 150 mg was the most cost- effective.
Poland,	CUA	NR	A vs. D D vs.	CHADS2-VASc = 1 and HAS- BLED = 0: NOK 882,000/QALY CHADS2-VASc = 2 and HAS- BLED = 1: NR	BLED = 1: NA CHADS2- VASc = 1 and HAS- BLED = 0: \notin 103444/QA LY ⁶ CHADS2- VASc = 2 and HAS- BLED = 1: NA NA	NR

Country , Year	Analysi s Time horizon	WTP	Compari son	Summary Metric	Summary Metric in 2019 Euros	Preferenti al Statement
2014	NR		warfarin			
			D vs. A, R	NR	NA	NR
Poland,	CUA Lifetim	NR	A vs. warfarin	NR	NA	NR
2013	e		A vs. aspirin	NR	NA	NR
			A vs. warfarin	£12,119/QALY	€15,795/QA LY ⁵	Apixaban was cost-
Scotland, 2013	CUA Lifetim e	NR	A vs. D	£13,467/QALY	€17,552/QA LY ⁵	effective compared with warfarin and dabigatran.
			A vs. R	NR	NA	Dominated
			E vs. warfarin	£23,539/QALY	€29,299/QA LY ⁵	Edoxaban dominated
			E vs. R	NR	NA	rivaroxaba
Scotland,	CUA	ND	E vs. D	NR	NA	n and was
2015	30 years	NR	E vs. A	NR	NA	as effective and less costly than apixaban.
Scotland, 2011	CUA Lifetim e	£20,000- 30,000/QAL Y (€26,067- 39,100/QAL Y)	D vs. warfarin	£6,986/QALY	€9,608/QAL Y ⁵	Dabigatran was more cost- effective than warfarin.
Scotland, 2012	CUA Lifetim e	£30,000/QA LY (€39,100/Q ALY)	R vs. warfarin	NR	NA	Dominated
Singapor e, 2018	CEA and CMA Lifetim e	NR	A, D, R vs. warfarin	<\$15,000/QALY	NA	NOACs were a cost- effective treatment option compared with

Country , Year	Analysi s Time horizon	WTP	Compari son	Summary Metric	Summary Metric in 2019 Euros	Preferenti al Statement
						warfarin for stroke prevention.
Spain, 2016	NA	NA	NA	NA	NA	NA
			Apixaban vs. warfarin	NR	NA	Apixaban and rivaroxaba
Sweden, 2013	CEA NR	NR	Apixaban vs. D, R	NR	NA	n were cost- effective compared with warfarin.
			E vs. warfarin	NR	NA	None of the
Sweden, 2016	CMA NR	NR	E vs. A, D, R	NR	NA	NOACs could be considered superior to others.

Abbreviations: A = apixaban; CEA = cost-effectiveness analysis; CMA = cost-minimization analysis; CUA = cost-utility analysis; D = dabigatran; E = edoxaban; HTA = health technology assessment; NA = not applicable; NOAC = non-vitamin K antagonist oral anticoagulant; NOK = Norwegian Krone; NR = not reported; QALY = quality-adjusted life year; R = rivaroxaban; VKA = vitamin K antagonist; WTP = willingness to pay

¹ Inflation rate at https://www.rba.gov.au/inflation/measures-cpi.html [15]

² Inflation rate at https://www150.statcan.gc.ca/t1/tbl1/en/tv.action?pid=1810000401 [16]

³ Inflation rate at https://www.banrep.gov.co/en/consumer-price-index [17]

⁴ Exchange rate at https://www.exchangerates.org.uk/EUR-COP-spot-exchange-rates-history-2019.html [18]

⁵ Inflation rate at https://ec.europa.eu/eurostat/web/hicp [19]

⁶ Inflation rate at https://www.ssb.no/en/kpi [20]

T	Number of Countries with			ntervention] e Following [
Interventio n	Positive Recommendatio ns	VKA	Apixaba n	Dabigatra n	Edoxaba n	Rivaroxaba n
Apixaban	Overall: 10/12* vs. VKAs: 8/12* vs. NOACs: 3/5*	8: Australia, Belgium, Canada, Ireland, Norway, Singapore, Sweden, England	NR	2: Ireland, Scotland	1: Canada	3: Canada, Ireland, Scotland
Dabigatran	Overall: 10/13 vs. VKAs: 8/13 vs. NOACs: 3/9	8: Australia, Belgium, Canada, Ireland, Netherland s, Norway, Singapore, England	1: Norway	NR	0	3: Norway, Australia, Sweden
Edoxaban	Overall: 5/7 vs. VKAs: 5/7 vs. NOACs: 2/5	5: Belgium, Canada, Scotland, England, Sweden	0	0	NR	2: Canada, Scotland
Rivaroxaba n	Overall: 10/13 vs. VKAs: 10/13 vs. NOACs: 2/8	10: Australia, Belgium, Canada, Ireland, Netherland s, Norway, Scotland, Singapore, Sweden, England	0	2: England, Netherlan ds	0	NR

Table 3. Summary of Cost-effectiveness and Recommendations Across NOAC vs. NOAC
and NOAC vs. VKA Comparisons by Country

Abbreviations: NOAC = non-vitamin K antagonist oral anticoagulant; NR = not reported; VKA = vitamin K

antagonist

*The HTA on apixaban from the Netherlands did not report economic results and was not included in the denominator.

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Table 4. Patient Access Information: Approval Dates, HTA Publication Date, and Match ofRecommendation to Approved Regulatory Label

Countr y	Apixaban		Dabigatran		Rivaroxaban		Edoxaban			Key
	Appro val	HT A Pu b	Appro val	HT A Pu b	Appro val	HT A Pu b	Appro val	HT A Pu b	Mat ch	Drivers or Conditio ns
Australi a	Jul 2011	Ma r 201 3 [*]	May 2011	Ma r 201 3 [*]	May 2012	Ma r 201 3 [*]	NA	NR	Yes	Cost
Colomb ia	Jul 2012	Ma y 201 6 ^x	Feb 2011	Dec 201 4 ^x	Feb 2012	Ma y 201 6 ^x	NA	NR	No	Rejected - cost
Netherl ands	Sept 2012	Feb 201 3 [^]	Aug 2013	Jun 201 2*	Jun 2012	Oct 201 2*	Feb 2015	Sep t 201 5 [^]	Yes	Clinical efficacy and safety
German y	Nov 2012	Jan 201 7 [*]	Aug 2011	Jan 201 7 [*]	Dec 2011	Jan 201 7 [*]	Jun 2015	Jan 201 7 [*]	NA	Clinical only
England	Nov 2012	Ma r 201 7*	Aug 2011	Ma r 201 2*	Dec 2011	Ma y 201 2*	Jun 2015	No v 201 7*	Yes	Clinical data and cost of INR monitori ng
Belgiu m	Nov 2012	Ma r 201 3 [°]	NA	NR	NA	NR	NA	NR	Yes	Clinical efficacy and safety
Ireland	Nov 2012	Ma y 201 3 [*]	Aug 2011	Au g 201 1 [*]	Dec 2011	Ma r 201 2 [*]	NA	NR	Yes	NR
Norway	Nov 2012	Ma r 201	Aug 2011	Ma r 201	Dec 2011	Ma r 201	NA	NR	Yes	Assumpti ons in model

	Apixaban		Dabigatran		Rivaroxaban		Edoxaban			Key
Countr y	Appro val	HT A Pu b	Appro val	HT A Pu b	Appro val	HT A Pu b	Appro val	HT A Pu b	Mat ch	Drivers or Conditio ns
		3*		3*		3*				
Poland	Nov 2012	Au g 201 3*	Aug 2011	Jun 201 4*	NA	NR	NA	NR	No	Price and narrow populatio n (CHADS 2 score ≥3 and ≥75 years old)
Scotlan d	Nov 2012	Jan 201 3 [*]	Aug 2011	Sep t 201 1*	Dec 2011	Feb 201 2 [*]	Jun 2015	Oct 201 5 [*]	Yes	Clinical efficacy and safety
Sweden	Nov 2012	Ma y 201 3 [*]	Aug 2011	Au g 201 6 [*]	Dec 2011	Au g 201 6 [*]	Jun 2015	Jun 201 6 [*]	Yes	Cost
Spain	Nov 2012	No v 201 6 [^]	NA	NR	NA	NR	NA	NR		NR
Canada	Dec 2012	Ma r 201 3 [*]	Oct 2010	Jul 201 3 [*]	Jan 2012	Jul 201 3 [*]	Oct 2016	Apr 201 7 [*]	Yes	Clinical data and cost
Brazil	Jul 2013	Feb 201 6	Dec 2011 ^x	Feb 201 6 ^x	Sept 2011	Feb 201 6 ^x	NA	NR	No	Rejected – Cost
Singapo re	Dec 2012	Oct 201 8 [*]	Mar 2011	Oct 201 8 ^x	Mar 2012	Oct 201 8 [*]	NA	NR	No	Rejected – Cost
France	NA	NR	NA	NR	NA	NR	Jun 2015	Jul 201	No	Clinical only –

	Apixaban		Dabigatran		Rivaroxaban		Edoxaban			Key
Countr y	Appro val	HT A Pu b	Appro val	HT A Pu b	Appro val	HT A Pu b	Appro val	HT A Pu b	Mat ch	Drivers or Conditio ns
							×	6^		heteroge neity in trial methods and populatio n

Abbreviations: HTA = health technology assessment; INR = international normalized ratio; NA = not applicable;

NR = not reported

* Recommended; ^ No recommendation; x Not recommended

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Highlights

- A review is needed of HTA submissions of clinical and/or economic value of NOACs.
- Cost-effectiveness was a major driver of recommendations based on WTP thresholds.
- Different thresholds limit ability to generalize recommendations across countries.