Cost-Effectiveness of Direct Non-Vitamin K Oral Anticoagulants Versus Vitamin K Antagonists for the Management of Patients with Non-Valvular Atrial Fibrillation Based on Available “Real-World” Evidence: The Italian National Health System Perspective

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Abstract
Background and objective The increasing availability of real-world evidence (RWE) about safety and effectiveness of direct non-vitamin K oral anticoagulants (DOACs) for the management of atrial fibrillation (AF) offers the opportunity to better understand the clinical and economic implications of DOACs versus vitamin K antagonists (VKAs). The objective of this study was to compare the economic implications of DOACs and VKAs using data from real-world evidence in patients with AF.

Methods A Markov model simulating the lifetime course of patients diagnosed with non-valvular AF was used to evaluate the cost-effectiveness of DOACs (i.e., rivaroxaban, dabigatran and apixaban) versus VKAs from the Italian National Health System (INHS) perspective. The model was made up of data from the literature and a meta-analysis of RWE on the incidence of stroke/systemic embolism (SE), major bleeding (MB), intracranial haemorrhage (ICH) and all-cause mortality (ACM); direct costs included drug costs, costs for drug monitoring, and management of events from official national lists. One-way and probabilistic sensitivity analyses (PSA) were used to assess the robustness of the results.

Results Results from the meta-analysis showed that apixaban had a high probability of being the most effective for stroke/SE, MB and ACM. Despite their higher acquisition costs, the cost-effectiveness analysis showed all DOACs involved a saving when compared with VKAs, with per-patient savings ranging between €4647 (rivaroxaban) to €6086 (apixaban). Moreover, all DOACs indicated a gain both in quality-adjusted life-years and life-years. According to PSA, findings related to apixaban were consistent, while for dabigatran and rivaroxaban PSA revealed a higher degree of uncertainty.

Conclusions The beneficial effect of DOACs on containing events showed in RWE had the potential to offset drug-related costs, thus improving the sustainability of treatment for non-valvular AF in daily clinical practice.

1 Introduction

Atrial fibrillation (AF) is the most common and clinically significant arrhythmia, and one of the major causes of stroke, heart failure, sudden death and cardiovascular morbidity in the world; it also carries a significant cost burden as a result of treatment and frequent hospitalization as well as considerable impairment in quality of life (QoL) [1, 2].

According to published data, about 33 million people suffer from AF [3]. A recent study also highlights wide variability in the prevalence of AF worldwide with significant gender differences. Specifically, estimates for 2010 suggested an overall (age-adjusted) prevalence of AF of about 6.0 (95% confidence interval (CI) 5.6–6.4) per 1000 among men and 3.7 (95% CI 3.5–4.0) per 1000 among women; these figures were slightly higher in industrialized countries (compared to developing countries) being, respectively, 6.6 (95% CI 6.0–7.4) per 1000 among men and 3.9 (95% CI 3.4–4.5) per 1000 among women. Moreover, the USA and Canada showed the highest prevalence, while central and northern Europe, as well as India, Japan and China had the lowest rates, with data from eastern countries likely to be underestimated [1, 3].
Sufficient evidence is now available to inform a cost-effectiveness analysis of direct non-vitamin K oral anticoagulants (DOACs) for the management of atrial fibrillation on the basis of real-word evidence.

Synthesizing available real-world evidence studies, apixaban, dabigatran and rivaroxaban were likely to improve health benefit over warfarin.

Despite the higher acquisition costs, apixaban was cost-effective compared to warfarin, suggesting savings for the Italian National Health System; considerable uncertainty still remained on the cost-effectiveness of dabigatran and rivaroxaban.

Several studies suggested an increasing trend in the prevalence and incidence of AF in the last decades; that course could partially be explained by the aging population and the increasing prevalence of co-morbidities and cardiovascular risk factors, in addition to other factors such as lifestyle changes and improved diagnosis [1, 3].

Management of patients with AF requires an integrated approach to monitor and control the disease, but also to prevent disease burden.

Stroke prevention is crucial in AF patients, and the use of oral anticoagulants has been demonstrated to reduce the risk of events and also to decrease mortality [4–6]; indeed, the 2016 European Society of Cardiology (ESC) guidelines for AF recommend the use of oral anticoagulants in all patients with CHA2DS2-VASC risk factors ≥ 2 [7].

At present vitamin K antagonists (VKAs; i.e., warfarin) and four diverse non-vitamin K oral anticoagulants (DOACs)—apixaban, dabigatran, rivaroxaban and edoxaban—are available on the market, and can be used for stroke prevention in non-valvular AF (NVAF).

While treatment with VKAs represented the standard for effective stroke prevention for many years, their use requires constant monitoring of the anticoagulation effect through measurement of the International Normalized Ratio (INR) to ensure an optimal level; this results in physical, psychological, social and financial consequences for the patient and the healthcare team [7, 8].

On the other hand, DOACs eliminate the need for laboratory monitoring, and have minor drug and food interactions and a wide therapeutic window [9]. As a drawback, without the need for monitoring, adherence to DOACs cannot be easily assessed.

Because of their ease of use in routine clinical practice and their excellent efficacy and safety profile [10], since their first introduction, DOACs have rapidly became the mainstay of therapy for stroke prevention in patients with NVAF [10].

Specifically, results from randomized controlled trials showed a significant reduction in the risk of mortality, bleeding and stroke for apixaban versus VKAs, while results for the other DAOCs are not as clear for both safety and efficacy endpoints [11–14]. However, regarding their relative safety and effectiveness in routine care, results have not been homogeneous among different studies and a branch of evidence has suggested increased risk for gastrointestinal bleeding related to the use of DOACs [15, 16].

Despite the number of publications and clinical trials conducted to evaluate the efficacy and safety of DOACs, not much effort has been applied to investigating the clinical and economic impact of the use of these drugs using real-world data and even trying to compare the different DOACs, particularly in Italy.

Indeed, evidence from “real-world” experiences may not perfectly match results from controlled settings, leading to discrepancies in both effectiveness and costs associated with the use of different therapies [17–19].

Accordingly, the primary objective of this study was to estimate the long-term cost-effectiveness of stroke prevention in patients with NVAF comparing available DOACs with the standard treatment, VKAs, taking into consideration the perspective of the Italian National Health System (INHS).

To this end, specific objectives of the present study were the collection and synthesis of available real-world evidence comparing DOACs versus VKAs for the prevention of stroke and management of NVAF patients; on the basis of that data an economic evaluation was performed adapting to the Italian context a previously developed model [20]. This study provides an evaluation of the cost-effectiveness of available direct oral anticoagulation therapies to assist clinicians and other healthcare decision-makers to make a more informed choice regarding patients’ treatment.

## 2 Methods

A systematic literature review followed by a network meta-analysis was performed to synthesize the available evidence related to the effectiveness and safety of the use of DOACs for the management of NVAF patients; results from the meta-analysis were then used as effectiveness inputs for the cost-effectiveness analysis.
2.1 Systematic Literature Review and Network Meta-Analysis

2.1.1 Systematic Review

The systematic literature review aimed at identifying available evidence about the effectiveness and safety of DOACs versus VKAs for the management of NVAF patients in a real-world setting and was performed according to the Preferred Reporting Items for Systemic Reviews and Meta-Analyses (PRISMA) statement method [21].

PubMed and Scopus databases were searched to retrieve studies published from 1 January 2009 to 31 September 2019 using the search strategies detailed in Online Supplementary Material (OSM) 1.

Studies published in scientific journals for which a full text was available in English and satisfying the inclusion and exclusion criteria reported in Table 1 were considered for inclusion in the systematic review.

In brief, retrieved studies were included if they referred to real-world studies comparing DOACs versus VKAs for the management of NVAF patients with CHA2DS2-VASC risk factors ≥ 2 and reported adjusted results (i.e., comparing propensity score-matched groups, multivariate models) related to the effect of the diverse treatments used at standard doses (apixaban 5 mg twice a day, dabigatran 150 mg twice a day, and rivaroxaban 20 mg once a day) on at least one of the following endpoints: stroke/systemic embolism (SE), major bleeding (MB), intracranial haemorrhage and all-cause mortality (ACM). Studies considering mixed populations treated with doses that were different from the standard were included if specific data for the different dosages were available, more than 50% of the population was treated at the standard dose, or if there was no clear evidence that the majority of the study population received doses different from the standard ones. Randomized trials or other experimental studies, studies considering specific subgroups of patients (i.e., NVAF patients with associated kidney disease, NVAF with CHA2DS2-VASC risk factors < 2), or those for which a standard and clear definition of events (i.e., referring to the specific ICD-9 or ICD-10 codes) was not available were excluded.

Two authors (SP and VL) independently performed the literature search, and screened retrieved studies for eligibility on the basis of titles (first step), abstracts (second step) and full text (third step). Any discrepancies were resolved involving a third author (GT) and reaching consensus.

The following data were obtained from studies included in the analysis: study characteristics, patients’ characteristics and outcomes; the number of events and population size or adjusted hazard ratio (HR) and its 95% CI were extracted and used in the network meta-analysis.

2.1.2 Network Meta-Analysis

A network meta-analysis was thus performed to combine and synthetize evidence from the systematic literature review. Network meta-analysis allows combining data from different studies assuming that within-studies estimates of relative treatment effects could be obtained as differences in the between-treatment effects, with the appropriate choice of a measurement scale.

Table 1 Summary of inclusion and exclusion criteria used in the systematic literature review

| Inclusion criteria | Exclusion criteria |
|--------------------|-------------------|
| **Type of studies** | Any observational studies (relying on real-world data): prospective cohorts, retrospective cohorts, case-control studies, registry analyses, analyses of administrative databases, etc. |
|                    | Randomized controlled trials or other experimental studies |
| **Population**     | Confirmed non-valvular atrial fibrillation (NVAF) |
|                    | Particular subgroups of patients with specific co-morbidities in addition to NVAF (i.e., renal disease, heart failure, etc.) |
| **Age ≥ 18 years** | |
| **Intervention**   | Direct oral anticoagulants (DOACs; alone or compared with warfarin) at standard doses (apixaban 5 mg twice a day, dabigatran 150 mg twice a day, rivaroxaban 20 mg once a day) |
|                    | Oral anticoagulant drugs excluding NAFV as the primary focus |
| **Outcomes**       | Any of the following: stroke (or stroke/systemic embolism), major bleeding, intracranial haemorrhage and all-cause death |
|                    | DOACs dosage clearly different from standards (in the majority of the study sample) |
The meta-analysis was performed considering the approach proposed by Woods et al. [22], which allows combining HR and cumulative count survival data on the log hazard scale and considering a Bayesian framework. The approach used also accounted for the correlations in relative treatment effect estimates that arise from trials with more than two treatment arms—multi-arm trials. Separated models were adapted for each endpoint of interest and analyses were performed using WinBUGS 1.4.3.

Each model was run for 50,000 burn-in simulations followed by an additional 200,000 runs. Two sets of initial values were used in all models and convergence was evaluated by trace plot; autocorrelation was also assessed. In addition, both fixed and random-effects models were considered for each endpoint of interest and the deviance information criteria (DIC) was used to compare the model fit. On the basis of the DIC value and on the estimates of the random-effect parameter, results from fixed models were retained for the analysis as there was no evidence concerning the significance of the random effect.

Results from the meta-analysis are expressed as HR and 95% credible interval (CrI); for each endpoint, ranking of the diverse alternatives was also derived from the model to provide a rank of the diverse treatment approaches.

2.2 Cost-Effectiveness Analysis

The cost-effectiveness analysis was performed on the basis of a Markov model considering a lifetime horizon and the INHS perspective. Results were reported using the incremental cost-utility ratio (ICUR) expressed as cost per quality-adjusted life-years (QALY) and the incremental cost-effectiveness ratio (ICER) expressed as cost per life-year gained (LYG). All costs and outcomes were discounted at 3.5% per year.

2.2.1 Model Structure and Perspectives

The economic evaluation was performed adapting a model previously developed as a Microsoft Excel enabled workbook; details about the structure of the model have been previously reported [20].

Adaptation of the previously developed model was performed to adjust it to the context of the INHS (a system providing universal coverage to citizens and residents, with public healthcare largely free of charge) and to account for the incidence of events from real-world studies.

The model evaluates and compares the use of DOACs for the prevention of stroke in AF and allows comparisons of each DOAC versus a VKA but also compares the cost-effectiveness of the diverse DOACs.

Markov transitions are employed to describe the disease progression, with a 3-monthly cycle.

A schematic representation of the model structure is provided in Fig. 1.

In brief the model mimics the disease course of a hypothetical cohort of 1000 patients diagnosed with NVAF. Patients entering the model because of the diagnosis of NVAF had predefined risks of events according to their actual condition [20, 23] that were modified according to the treatment arm they entered in. Accordingly, from the initial state “AF well” patients move to different states characterized by the number of events they experience—“AF + single event”, “AF + two events”, “AF + three events” or even death. Stroke/SE, major bleed (MB) or intracranial haemorrhage (ICH) were considered as events, and transition probabilities were defined on the basis of the number and type of events experienced.

Four different treatment arms were considered in the analysis, representing first-line treatment with warfarin, apixaban, dabigatran and rivaroxaban. Edoxaban was not considered in the present analysis as just one study was retrieved from the literature search performed.

The following assumptions were also considered in the model: (1) patients always switch to no treatment after haemorrhage; (2) patients may switch from DOACs to warfarin or from warfarin to no treatment after stroke, major bleed or systemic embolism; (3) patients may switch from DOACs to warfarin or discontinue warfarin at any time due to lack of compliance.

The model estimates annual direct health costs for the different treatment options considering costs associated with oral anticoagulant therapy, costs related to hospitalizations for the management of events and monitoring costs, where appropriate.

2.2.2 Effectiveness and Cost Data

Inputs for effectiveness were derived from the meta-analysis on real-world experience with the use of DOACs versus VKAs described above. On the basis of mortality and the risk of events, life-years (LYs) and QALYs for the diverse treatments were thus obtained from the model. To derive QALYs, utilities were applied according to previously

![Fig. 1 Structure of the Markov model used for the cost-effectiveness analysis](image-url)
published data referring to the events of interests and considering a utility value for stable AF equal to 0.779 [24–26]; details of the values used in the analysis are reported in Table 2.

Direct health costs related to the oral anticoagulant therapy (either VKAs and DOACs), direct health costs of events and eventual monitoring costs were considered in the analysis.

Specifically, annual cost associated with the oral anticoagulant therapy was estimated multiplying, for the different treatment options, daily standard dosage times unit costs of the product considering ex-factory prices from the National Agency for Drugs [27] and from the National Gazette [28–30], including statutory discount. For VKAs, costs of monitoring were also added to the cost of drugs to fully capture all treatment costs; costs of INR monitoring in the Italian setting were derived from previously published data and inflated to the reference year [31, 32].

Costs associated with the incidence of events were estimated considering annual incidence rate and reimbursement associated with hospitalization for the management of each event according to official national charges set by the Italian Ministry of Health (i.e., Tariffario prestazioni assistenza ospedaliera per acuti) [33]. Only for stroke costs associated with the event also included costs of rehabilitation derived from a previous study conducted in Italy [34] and inflated to the reference year. Details of cost inputs used are reported in Table 3.

All costs are expressed in Euros and referred to 2019 values.

2.2.3 Sensitivity Analysis

To evaluate the robustness of results both one-way (OWSA) and probabilistic sensitivity analyses were conducted. One-way sensitivity analysis was conducted on all model parameters associated with uncertainty, modifying base-case inputs by ± 20% and showing main results from the analysis in a Tornado diagram. In the probabilistic sensitivity analysis (PSA), costs, utilities and transition probabilities were varied simultaneously over their 95% CIs (see OSM 2) and ICURs and ICERs were calculated over 1000 simulations. Results from the PSA were presented in a cost-effectiveness plane.

### Table 2 Disutility values used in the base-case analysis

| Events                        | Disutility value |
|-------------------------------|------------------|
| Stroke                        | −0.590           |
| Major bleeding                | −0.030           |
| Intracranial haemorrhage      | −0.179           |
| Systemic embolism             | −0.131           |

### Table 3 Details of cost inputs used in the base-case analysis

| Resource type                        | Unit costs | Costs/day |
|--------------------------------------|------------|-----------|
| Stroke (DRG 14+rehabilitation)       | €19,624    | –         |
| Major bleeding (DRG 174, 175, 14)    | €3568*     | –         |
| Intracranial haemorrhage (DRG 14)   | €3891      | –         |
| Transient ischaemic attack           | €2967      | –         |
| Apixaban                             | €0.212/mg  | €2.12     |
| Dabigatran                           | €0.007/mg  | €2.12     |
| Rivaroxaban                          | €0.104/mg  | €2.11     |
| Warfarin                             | €0.01€     | –         |

*This value was obtained as the weighted average of the tariff associated with DRG 174, 175, 14 where the number of hospital admissions from the Italian Ministry of Health was used as weight.

3 Results

3.1 Systematic Literature Review and Network Meta-Analysis

Articles identified through the literature searches were checked to identify any duplicates. Then reviewers screened titles and abstracts and selected publications that met the eligibility criteria; fulltexts of selected references were then assessed.

A total of 581 references were identified through the searching strategy, 494 were identified in PubMed, 84 from Scopus and three additional studies were added as being previously known.

After excluding duplicates, a total of 568 references were independently screened and according to pre-defined inclusion and exclusion criteria 424 studies were excluded on the basis of title or abstract. Finally, 144 full texts were evaluated, and among these overall 42 studies were included in the meta-analysis. Figure 2 provides the flow diagram followed in our review to select the retrieved studies.

Data for each outcome were extracted and organized by the standard dosages and evaluated in the meta-analysis.

Details of studies included the meta-analysis and a summary of main details in terms of study type, interventions considered and the type of endpoints analysed are reported in OSM 3.

Accordingly 26 studies were considered for stroke/SE [16, 35–60], 29 for MB [16, 36, 40–44, 46–48, 50, 52, 54–56, 58, 61–72], 20 for ICH [36, 38, 41–43, 45, 49–52, 54, 55, 60, 63, 64, 71–76] and 13 for ACM [16, 36, 41–43, 46, 52–54, 67, 71, 72, 77, 78].

Results from the meta-analysis suggested that for each endpoint all DOACs significantly reduced the risk of events and mortality; only when considering major bleeding the network meta-analysis suggested that rivaroxaban had a
higher probability of implying a slight increased risk as compared to VKAs (Table 4).

Trying to rank the effectiveness and safety profile of the different treatments with respect to events considered in the analysis, apixaban resulted in having a high probability of being the best option for both stroke/SE, MB and ACM. On the other hand, for ICH dabigatran showed a higher probability of being the best alternative (Table 5).

### 3.2 Cost-Effectiveness Analysis

Despite the lower acquisition costs of the drug, overall lifetime direct costs estimated through the model were higher for warfarin, €21,331 per patient, while apixaban resulted in the lowest one, €15,245 per patient. Costs associated with dabigatran and rivaroxaban were slightly higher than those associated with apixaban, €16,003 and €16,684 per patient, respectively (Table 6).

As shown in Fig. 3, the overall costs associated with warfarin were mainly driven by management costs that also comprised costs for INR monitoring; the rest were almost all event costs. On the contrary, costs associated with DOACs were for the large part equally driven by events costs and drug costs.

| Table 4 Effectiveness inputs from the network meta-analysis |
|---------------------------------|-----------------|
| **Drugs**                       | **HR (95% CrI)** |
| **Stroke/systemic embolism**    |                 |
| Apixaban vs. warfarin           | 0.755 (0.718–0.796) |
| Dabigatran vs. warfarin         | 0.887 (0.837–0.938) |
| Rivaroxaban vs. warfarin        | 0.857 (0.821–0.894) |
| **Major bleeding**              |                 |
| Apixaban vs. warfarin           | 0.594 (0.576–0.613) |
| Dabigatran vs. warfarin         | 0.741 (0.716–0.766) |
| Rivaroxaban vs. warfarin        | 1.034 (1.010–1.058) |
| **Intracranial haemorrhage**    |                 |
| Apixaban vs. warfarin           | 0.601 (0.553–0.654) |
| Dabigatran vs. warfarin         | 0.484 (0.432–0.540) |
| Rivaroxaban vs. warfarin        | 0.710 (0.667–0.763) |
| **All-cause mortality**         |                 |
| Apixaban vs. warfarin           | 0.706 (0.676–0.737) |
| Dabigatran vs. warfarin         | 0.750 (0.716–0.785) |
| Rivaroxaban vs. warfarin        | 0.883 (0.858–0.909) |

**CrI** credible interval

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**Fig. 2 Preferred reporting items for systemic reviews and meta-analyses (PRISMA) flow diagram showing results of the search strategy**

- Records identified through PUBMED searching (n = 494)
- Records identified through SCOPUS searching (n = 84)
- Additional records identified (n = 3)
- Records after duplicates removed (n = 568)
- Records screened (n = 568)
- Records excluded After Title/Abstract Screening (n = 424)
- Full-text articles assessed for eligibility (n = 144)
- Full-text articles excluded, with reasons (n = 102)
- Studies included in quantitative synthesis (meta-analysis) (n = 42)
The cost-effectiveness analysis showed that all DOACs lead to savings when compared with VKAs, resulting in a reduction of per patient cost ranging between €4647 (rivaroxaban) and €6086 (apixaban). Dabigatran incurred a reduction in per patient costs of about €5327 when compared with VKAs. Moreover, all DOACs implied a gain in both QALYs and LYs.

Accordingly, when compared to VKAs all DOACs resulted in dominantly negative values for ICURs and ICERs, induced by lower costs and higher effectiveness (Table 6).

On the other hand, comparisons among the different DOACs highlighted that apixaban had the lowest per-patient costs, with QALYs and LYs that were slightly higher than those seen with the other DOACs (Table 7).

As compared to apixaban, rivaroxaban and dabigatran resulted in higher costs and lower effectiveness (in term of both LYs and QALYs).

The main results from the OWSA are shown in Fig. 4. When comparing DOACs and VKAs, for all DOACs drug costs were among the main drivers of the analysis; variation in the HR for ACM also had a sensible impact on results from the base-case analysis, particularly when comparing apixaban and dabigatran versus VKAs. Results from the PSA suggested that results obtained from the base-case analysis showed a high degree of uncertainty with regard

### Table 5 Probability of treatment ranking for the different events considered

| Events                          | Best  | Second best | Third best | Fourth best |
|---------------------------------|-------|-------------|-----------|-------------|
| Stroke/systemic embolism        |       |             |           |             |
| Warfarin                        | 0.00  | 0.00        | 0.00      | 1.00        |
| Apixaban                        | 1.00  | 0.00        | 0.00      | 0.00        |
| Dabigatran                      | 0.00  | 0.11        | 0.89      | 0.00        |
| Rivaroxaban                     | 0.00  | 0.89        | 0.11      | 0.00        |
| Major bleeding                  |       |             |           |             |
| Warfarin                        | 0.00  | 0.00        | 1.00      | 0.00        |
| Apixaban                        | 1.00  | 0.00        | 0.00      | 0.00        |
| Dabigatran                      | 0.00  | 1.00        | 0.00      | 0.00        |
| Rivaroxaban                     | 0.00  | 0.00        | 1.00      | 0.00        |
| Intracranial haemorrage         |       |             |           |             |
| Warfarin                        | 0.00  | 0.00        | 0.00      | 1.00        |
| Apixaban                        | 0.00  | 1.00        | 0.00      | 0.00        |
| Dabigatran                      | 1.00  | 0.00        | 0.00      | 0.00        |
| Rivaroxaban                     | 0.00  | 0.00        | 1.00      | 0.00        |
| All-cause mortality             |       |             |           |             |
| Warfarin                        | 0.00  | 0.00        | 0.00      | 1.00        |
| Apixaban                        | 0.98  | 0.02        | 0.00      | 0.00        |
| Dabigatran                      | 0.02  | 0.98        | 0.00      | 0.00        |
| Rivaroxaban                     | 0.00  | 0.00        | 1.00      | 0.00        |

### Table 6 Results from the cost-effectiveness analysis for direct oral anticoagulants compared to vitamin K antagonists

| Drugs     | Overall costs | QALYs | LYs  | Δ costs | Δ QALYs | Δ LYs | ICER (€/QALYs) | ICER (€/LYs) |
|-----------|---------------|-------|------|--------|--------|-------|----------------|--------------|
| Warfarin  | €21,331       | 5.09  | 8.86 | –      | –      | –     | –              | –            |
| Apixaban  | €15,245       | 5.90  | 10.18| – €6086| 0.81   | 1.33  | – €7489        | – €4593      |
| Dabigatran| €16,003       | 5.77  | 9.92 | – €5327| 0.66   | 1.06  | – €8085        | – €5006      |
| Rivaroxaban| €16,684      | 5.39  | 9.31 | – €4647| 0.30   | 0.46  | – €15,423      | – €10,208    |

LYs: life-years, QALYs: quality-adjusted life-years, ICER: incremental cost effectiveness ratio

### Fig. 3 Results from the cost-effectiveness analysis: per-patient costs according to treatment arm
to the results from the cost-effectiveness analysis related to the comparison of dabigatran and rivaroxaban versus VKAs, while the scatter cloud related to apixaban was almost consistently located in the second quadrant, suggesting robustness of the findings from the base-case analysis (Fig. 5).

4 Discussion

To our knowledge this is the first study conducted to assess and compare the cost-effectiveness of the use of different DOACs versus VKAs considering a real-world setting and the Italian context.

Excluding edoxaban, which has been launched more recently, and for which just a few studies were available at the time of the analysis, with just one meeting the inclusion and exclusion criteria, at present there is sufficient real-world evidence to support the use of other DOACs in terms of both effectiveness and costs.

Results from the present study highlight a clear benefit related to the use of DOACs versus VKAs because of the higher QALYs and LYs induced by their effectiveness and safety profile and also their lower costs; indeed, the higher acquisition costs of DOACs were offset by reduced costs associated with management of events and patient monitoring. Specifically, on the basis of the analyses performed, apixaban resulted in higher benefits as compared to the other DOACs.

Results from the cost-effectiveness analysis also suggest the potential cost-effectiveness of all DOACs, although PSA clearly indicated that only for apixaban results were consistent, while a high degree of uncertainty was noted for both dabigatran and rivaroxaban.

Few other studies have been conducted focusing on the cost-effectiveness of DAOCs in a real-world setting [18, 79], while a consistent branch of evidence deals with the cost-effectiveness of these drugs considering data from RCTs [80–84].

Informing decision-making evidence from real-world settings is essential, even if not required from regulatory bodies, to adjust the relative efficacy results from RCTs to the actual performance of the comparator, to better contextualize the analysis to specific settings (i.e., in terms of patient characteristics, treatment scenarios), and to expand findings beyond the trial duration [18].

Findings from the present study are somewhat in line with a recent cost-effectiveness analysis on real-world evidence aimed at evaluating the use of available DOACs (vs. VKAs) considering the perspective of the Dutch National Health System [79].

As real-world practice may largely vary across different contexts, a limitation of the present study is that the analysis performed is based on real-world data from diverse contexts. Despite inclusion and exclusion criteria used for the identification of evidence included in the meta-analysis were defined also to ensure the inclusion of studies being as similar as possible (in terms of study design, patients, etc.), being real-world studies reporting evidence from clinical practice in different contexts a certain degree of homogeneity in the evidence cannot be excluded. Moreover, not enough data

Table 7 Results from the cost-effectiveness analysis: difference in per patient costs, QALYs and LYs between the different direct oral anticoagulants

| Drugs   | Costs difference (Euro) | Apixaban | Dabigatan | Rivaroxaban |
|---------|-------------------------|----------|-----------|-------------|
| Apixaban| –                       | –        | –         | –           |
| Dabigatan| €758.53                | –        | –         | –           |
| Rivaroxaban| €1438.65             | €680.11  | –         | –           |

| Difference in QALYs |
|---------------------|
| Apixaban            | Dabigatan | Rivaroxaban |
| Apixaban            | –        | –          |
| Dabigatan           | − 0.15   | –          |
| Rivaroxaban         | − 0.51   | − 0.36     |

| Difference in LYs  |
|-------------------|
| Apixaban          | Dabigatan | Rivaroxaban |
| Apixaban          | –        | –          |
| Dabigatan         | − 0.26   | –          |
| Rivaroxaban       | − 0.87   | − 0.61     |

LYs life-years, QALYs quality-adjusted life-years
Real-World Cost-Effectiveness of DOACs

The real-world setting could strongly impact the effectiveness and safety of drugs, and thus also on events rate and costs. Moreover, as only few real-world studies reported data on the impact of the diverse treatments on both myocardial infarction (MI) and transient ischemic attack (TIA), these events were not considered in the model. Despite this result, some specific insights into the Italian context are available at present to perform a comprehensive cost-effectiveness analysis based on robust evidence in the Italian context. Future studies, when feasible, could offer more specific insight into the Italian context.

Another study limitation lies in the fact that the model did not specifically account for adherence with the different DOACs as limited and inconsistent data were found from the systematic literature review. Indeed, despite it being unfeasible to model the impact of adherence on drug-related costs and on other direct health costs, being based on real-world studies, we believe the evidence used in the meta-analysis performed allow us to take into account the possible effect of suboptimal adherence behaviour on the incidence of events.

This point merits further study as adherence in a real-world setting could strongly impact the effectiveness and utility ratio, MB major bleeding, SE systemic embolism, LYs life-years, QALYs quality-adjusted life-years

Fig. 4 Tornado diagrams showing the main results from the one-way sensitivity analysis (OWSA). ICH intracranial haemorrhage, ICER incremental cost effectiveness ratio, ICUR incremental cost utility ratio.
meta-analysis and cost-effectiveness evaluation performed focusing on RCTs, edoxaban was shown to have a favourable effectiveness profile [23], resulting in being ranked as the second best after apixaban, in view of its effect related to the risk reduction for both MB and mortality versus VKA.

Despite the limitations presented, the present study adds to the limited evidence produced on the cost-effectiveness of DOACs in the real-world, contributing to the removal of uncertainties about DOACs [84] and also providing useful data for both clinicians and decision makers in Italy not only on the use of DOACs compared to VKAs, but also providing suggestions for choosing the most effective and sustainable options among actually available alternatives.

5 Conclusions

Based on up-to-date evidence from real-world studies, the present analysis suggests that apixaban is a cost-saving alternative to warfarin therapy for the prevention of stroke in patients with AF in the Italian healthcare setting; however, considerable uncertainty still remains on the cost-effectiveness of dabigatran and rivaroxaban.

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Fig. 5 Cost-effectiveness plane resulting from the probabilistic sensitivity analysis (PSA). QALYs quality-adjusted life-years, WTP willingness-to-pay

Declarations

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Consent to participate Not applicable.

Consent for publication Not applicable.

Data availability The data used in this study are fully available in databases.

Code availability Not applicable.

Conflict of interest VL, SP and GT are employees of the Institute of Management, Scuola Superiore Sant’Anna, which received financial support from Pfizer and Bristol Myers Squibb in connection with the development of this article. The authors have no other conflicts of interest to declare.

Author contributions VL conceived the paper and performed the analysis. SP performed the literature search, all the authors were involved in the systematic literature review. All authors were involved in the study design, discussion and interpretation of the results. All authors repeatedly edited the manuscript and approved the final version.

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