Cost-Effectiveness Analysis of Apixaban Versus Edoxaban in Patients with Atrial Fibrillation for Stroke Prevention

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Abstract

Objective Our objective was to assess the cost effectiveness of apixaban versus edoxaban in the prevention of stroke and systemic embolism (SE) in patients with atrial fibrillation (AF) in Spain.

Methods We customized a Markov model with ten health states to estimate the lifetime economic and clinical outcomes in 6-week cycles. The efficacy (clinical event rates per 100 patient-years) and safety data were derived from a pairwise indirect treatment comparison. The analysis was conducted from both the national health service (NHS) and societal perspectives, and included pharmaceutical costs (retail price plus value-added tax (VAT) and applicable national deductions) according to daily dosages (apixaban 10 mg (5 mg twice daily (bid)) and edoxaban 60 or 30 mg) and complications and disease-management costs, obtained from national databases. Utilities for quality-adjusted life-year (QALY) calculations reflected EuroQoL 5-Dimension scores in patients with AF. An annual discount rate of 3% was applied for costs (€, year 2019 values) and outcomes.

Results In a 1000-patient cohort, apixaban 5 mg bid versus edoxaban 60 mg could avoid five strokes, six major bleedings and 29 clinically relevant non-major bleedings (CRNMBs). Compared with edoxaban 30 mg, apixaban could avoid 21 strokes and two SEs. An increase in bleedings was observed with apixaban (seven haemorrhagic strokes, 48 major bleedings and 17 CRNMBs). Apixaban yielded 0.04 additional QALYs compared with edoxaban 60 mg or 30 mg. Incremental costs/QALY were €9639.33 and €354.22 for apixaban versus edoxaban 60 mg and edoxaban 30 mg, respectively, from the NHS perspective and €7756.62 for apixaban versus edoxaban 60 mg from the societal perspective. Apixaban was dominant versus edoxaban 30 mg from the societal perspective. Sensitivity analyses confirmed the robustness of the model.

Conclusions This study suggests that apixaban 5 mg bid is a cost-effective alternative to edoxaban for stroke prevention in the AF population in Spain.

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Key Points for Decision Makers

Apixaban (5 mg twice daily) is an effective therapy for the treatment of atrial fibrillation (AF), preventing events such as strokes and systemic embolism compared with edoxaban.

Costs per patient treated with apixaban are similar to those in patients receiving edoxaban 60 mg or 30 mg daily.

Apixaban could be considered a cost-effective option versus edoxaban for stroke prevention in patients with AF in Spain.
Atrial fibrillation (AF) is the most frequent cause of cardiac arrhythmia and is among the main causes of stroke and thromboembolic events [1]. It is associated with increased morbidity and mortality and high medical costs related to the substantial consumption of health resources for its management [2]. In Spain, the cost to the health system is up to €13,319 per patient with cardioembolic ischaemic stroke [3]. The total cost in patients admitted to stroke units equals €27,711 per patient, with up to €18,643 related to non-healthcare costs [4].

Although the current European clinical guidelines for patients with AF recommend anticoagulation therapy as a preventive measure for associated complications [5], more than 40% of affected Spanish patients are not treated according to these recommendations [6].

Traditionally, the elective treatment for stroke prevention in patients with AF has been based on the administration of a vitamin K antagonist (VKA), mainly acenocoumarol, with the consequent risk of developing treatment-related haemorrhage [7].

The discovery some years ago of direct-acting oral anticoagulants was a significant development. This drug class includes apixaban, dabigatran, rivaroxaban and, most recently, edoxaban. Their main advantage is that they do not require the continuous and permanent monitoring of international normalized ratio (INR) and dose adjustment to assure therapeutic drug concentrations that are required for VKAs.

The efficacy and safety of the direct-acting anticoagulant agents for stroke prevention in patients with AF have been demonstrated in many clinical trials: ARISTOTLE [8] (apixaban 5 mg twice daily (bid) vs. warfarin), AVERROES [apixaban vs. acetylsalicylic acid (ASA)] [9], RELY (dabigatran vs. warfarin) [10], ROCKET-AF (rivaroxaban vs. warfarin) [11] and ENGAGE-AF (edoxaban vs. warfarin) [12].

Several economic evaluations have provided additional information to the previous efficacy evidence for health decision makers. The efficiency of these drugs has been assessed in different settings and with different comparators. These studies include cost-effectiveness analyses of apixaban versus warfarin, acenocoumarol, ASA, dabigatran and rivaroxaban [13] and, most recently, a cost-utility analysis versus edoxaban in the UK [14]. Evaluations specifically for the Spanish setting include those for apixaban versus acenocoumarol [15], apixaban versus dabigatran [16] and apixaban versus rivaroxaban [17].

Since edoxaban, a new therapeutic option, is now available in Spain, this study aimed to assess the cost effectiveness of apixaban versus edoxaban for the prevention of stroke and systemic embolism (SE) in patients with AF in Spain to provide additional information for the medical decision-making process.

### 2 Material and Methods

A previously developed and validated Markov model [18, 19], simulating the evolution of AF, was customized to the Spanish setting. A panel of Spanish experts, comprising three clinicians (two cardiologists and one internist) and five health economists, who are among the authors of the present manuscript, validated the representativeness of the parameters used and provided information about the local management of patients. Clinicians completed online questionnaires, and a face-to-face meeting was held with all the experts and authors for discussion and to reach consensus related to the final input values.

A hypothetical cohort of 1000 patients eligible to be treated with apixaban or edoxaban was assessed to show potential events avoided in understandable figures. The clinical characteristics of this cohort, including average age (70 years), the proportion of females (35.3%) and CHADS2 average score (2.1), were defined based on the population included in the ARISTOTLE clinical trial [8], which assessed the efficacy and safety of apixaban and warfarin and were considered representative of Spanish patients with AF by the expert panel.

The model used for the present analysis represents the disease course, including ten main mutually exclusive health states: AF, ischaemic stroke, haemorrhagic stroke, other intracranial haemorrhages, other major bleedings, clinically relevant non-major bleedings, myocardial infarction (MI), SE, treatment discontinuation and death related to any cause different to the previously described clinical events. Based on the modified Rankin scale (mRS), the ischaemic or haemorrhagic strokes were categorized as mild (mRS 0–2), moderate (mRS 3–4) or severe (mRS 5).

The simulation was initiated with the whole cohort in the AF state. Throughout the simulation, to reflect a potential real-life patient evolution, the patients could remain in or transition through the previously described health states (Fig. 1). Patients remained in the AF state until stroke, bleeding, SE, MI, treatment discontinuation or death occurred.

Transitions between the discrete health states were allowed in 6-week cycles (deliberately defined to capture the possibility of events related to AF occurring within such a short timeframe), with probabilities based on the likelihood of the different clinical events depending on the assigned treatment, reflecting efficacy and safety patterns for the assessed therapies.

Like the death state, the states of SE, ischaemic stroke and haemorrhagic stroke (mild, moderate and severe) and MI were permanent states, with the rest being transient health
states occurring for a maximum of 6 weeks before returning to the prior or moving to a subsequent health state.

A half-cycle correction approach was applied [20].

Hospitalizations for cardiovascular reasons other than those related to MIs or strokes were modelled in the background and not as health states.

2.1 Alternatives

This analysis is the last of a series of economic evaluations of apixaban compared with other anticoagulant treatments, such as acenocoumarol and other direct-acting oral anticoagulants (dabigatran and rivaroxaban), that were previously authorized and launched.

The present analysis focused on assessing apixaban compared with edoxaban, the latest direct-acting oral anticoagulant to become available on the Spanish market.

Given the clinical data reported from the ENGAGE-AF clinical trial [12], high and low doses of edoxaban were assessed in the present model (60 or 30 mg daily, which is the recommended dose in patients with AF with moderate or severe renal impairment, low body weight ≤ 60 kg and/or concomitant use of cyclosporin, dronedarone, erythromycin or ketoconazole) [21].

The assessed dosage of apixaban was 5 mg bid [22], consistent with the dosage received by most of the patients in the ARISTOTLE trial [8] (< 4.7% of patients received the apixaban low dose, recommended for those aged ≥ 80 years or with body weight ≤ 60 kg or serum creatinine ≥ 1.5 mg/dL) and given that the published efficacy was reported for patients mainly treated with this dosage.

Aligned with the assumption used in previous customizations of the same model [14–19] and validated by the expert panel, the administration of ASA was considered a...
second-line treatment in patients who stopped or withdrew from the first-line therapy (any of the two drugs assessed).

### 2.2 Clinical Events

In the absence of head-to-head clinical trials, the clinical event rates per 100 patient-years were derived from a pairwise indirect treatment comparison analysis (detailed information can be found in a previous publication [14]), which was performed using the Bucher method [23]. In the indirect comparison, the relative hazard ratios (HRs) were estimated in relation to the existing common comparator, warfarin, from the apixaban and edoxaban clinical trials (ARISTOTLE [8] and ENGAGE-AF [12], respectively).

As in the previously published economic evaluation of apixaban versus edoxaban [14], the event rates and the HR (summarized in Table 1) were estimated with the rates observed per CHADS2 score level (CHADS2 0–1 (34.0%), CHADS2 2 (35.8%), CHADS2 ≥ 3 (30.2%)) and INR control based on the centre’s median time in therapeutic range.

For ASA, the event rates were derived from a subgroup of patients with prior VKA exposure from the AVERROES trial [9].

Case-fatality rates following the occurrence of each event derived from trials were applied. All-cause mortality excluding deaths attributable to stroke, bleeding, MI and SE were applied for patients in the AF health state to avoid double counting. Beyond 1.8 years (consistent with trial duration), mortality was modelled based on age- and sex-specific general mortality. The general mortality for males and females aged < and > 75 years were derived from published Spanish mortality data [24], which were modelled fitting a Gompertz survival function (details about parameters for the distribution function are available on request). The use of the Gompertz function instead of the raw data allowed a more refined estimation of the risk of mortality for every 6-week cycle (compared with yearly data on general life tables). In addition to background mortality, the model implemented adjustment factors, such as the HR, to reflect the potential increase in mortality rates associated with clinical events (stroke, MI, SE and bleedings). Since mortality due to strokes, MI, SE and bleedings was explicitly modelled at the occurrence of the event, increased mortality for patients with AF due to these causes was excluded from the calculation of the HR to avoid double counting.

Survival outcomes, estimated as life-years gained (LYG), were used to calculate quality-adjusted life-years (QALYs) by means of the utility values representing the patients’ preference for certain health states. In the present model, a basal utility associated with AF (0.727) and different utilities for the health states were used. Temporal decrements of utility in cases of complications were also applied (Table 2). All values for utilities and decrements were consistent with those previously used in international [14, 18, 19] and national [15–17, 25] publications. These data are referred to as the scores of the EuroQol 5-Dimensions (EQ-5D) questionnaire obtained in a sample of patients in the UK [26]. Further details about modelling can be found in previous publications [14, 18, 19], and additional information is also available on request.

### 2.3 Perspective, Discount Rate and Time Horizon

The analysis was carried out from two perspectives: that of the national health service (NHS) and the societal perspective.

The NHS perspective included direct medical costs related to drug acquisition, complication management, monitoring and AF patient follow-up. The societal perspective also included non-medical costs, but productivity costs were excluded due to lack of reliable data.

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**Table 1 Clinical event rates**

| Event                                      | Average rate per 100 patient-years (95% CI) | HR vs. apixaban (95% CI) |
|--------------------------------------------|-------------------------------------------|--------------------------|
|                                            | Apixaban ASA as subsequent treatment       | Edoxaban 30 mg Edoxaban 60 mg |
| Ischaemic stroke                           | 0.98 (0.56–1.52) 3.45 (1.97–5.34)          | 1.48 (1.12–1.96) 1.04 (0.78–1.39) |
| Intracranial haemorrhage                   | 0.33 (0.19–0.51) 0.32 (0.18–0.50)          | 0.74 (0.46–1.20) 1.11 (0.71–1.73) |
| Major bleedings                            | 1.79 (1.02–2.77) 0.89 (0.51–1.37)          | 0.66 (0.53–0.83) 1.13 (0.91–1.39) |
| Clinically relevant non-major bleedings    | 2.08 (1.19–3.22) 2.94 (1.68–4.54)          | 0.95 (0.80–1.13) 1.25 (1.06–1.48) |
| Other anticoagulant treatment discontinuations | 13.18 (7.53–20.38) NA                      | 1.04 (0.96–1.13) 1.10 (1.01–1.19) |
| Myocardial infarction                      | 0.53 (0.30–0.82) 1.11 (0.63–1.72)          | 1.37 (0.95–1.96) 1.07 (0.74–1.55) |
| Systemic embolism                          | 0.09 (0.05–0.14) 0.40 (0.23–0.63)          | 1.39 (0.57–3.36) 0.74 (0.29–1.92) |
| Other cardiovascular hospital admissions   | 10.46 (5.98–16.17) 13.57 (7.76–20.98)      | 1.00 (0.90–1.10) 1.00 (0.90–1.10) |
| Mortality by other causes                  | 3.08 (2.50–3.72) NA                        | 1.00 (0.90–1.10) 1.00 (0.90–1.10) |

ASA acetylsalicylic acid, CI confidence interval, HR hazard ratio, NA not applicable

*Mortality related to any cause, excluding stroke, haemorrhage, myocardial infarction and systemic embolism*
A 3% annual discount rate was applied to both costs and health outcomes, following the latest published recommendations for the development of economic evaluations in Spain [27].

A lifetime period was chosen for the time horizon. Coincident with previously published evaluations of apixaban compared with other alternatives performed in Spain [16, 17], this represents an average life expectancy of 80.4 years, which was considered by the expert panel to be applicable to patients with AF.

### 2.4 Resource Consumption and Costs

The estimation of total costs included the following direct medical costs: anticoagulant therapy acquisition, complication management in acute and maintenance phases, dyspepsia related to anticoagulant treatment, renal monitoring and clinical follow-up for AF. The acute phase, based on experts’ validation, was assumed to last 2 weeks, involving the time for hospital care and rehabilitation. Following this acute care period, maintenance phase costs were accumulated until the patient’s death.

The pharmaceutical costs for each of the two drugs assessed were calculated based on the daily dosages (apixaban 5 mg bid, edoxaban 30 mg and edoxaban 60 mg) stated on the summaries of product characteristics [21, 22] and the public retail prices [28] plus a VAT (4%), with the applicable deduction established by Royal Decree-Law 8/2010 [29].

The management costs of complications during the acute phase were derived from the average costs of diagnostic-related group official prices established by the autonomous regions [30]. The management costs of complications in maintenance, estimated as monthly costs, were obtained from several published Spanish sources [31–35].

The cost of dyspepsia related to anticoagulant treatment was included as a monthly clinical management cost obtained from the literature [36], considering the frequency (1.67%) observed in the ARISTOTLE trial [8] for apixaban and assuming an equivalent value for edoxaban. The renal monitoring cost was estimated as an annual cost, considering a yearly determination for each of the assessed therapies [37]. The cost of AF clinical follow-up was based on local clinical practice and validated by the expert panel, with a routine visit every 3 months assumed, regardless of the treatment chosen. Non-medical costs were obtained from the Spanish literature for both acute and maintenance costs [38], including societal costs related to the management of patient dependency, such as housing adaptation and informal care.

All costs are expressed in € (year 2019 values). When required, the consumer price index provided by the Statistic National Institute was applied. The unit costs used in the model are detailed in Table 3.

### 2.5 Cost-Effectiveness Analysis

The incremental cost-effectiveness ratio (ICER) in terms of cost per additional LYG and the incremental cost-utility ratio (ICUR) in terms of cost per additional QALY were estimated by means of differences in costs and health outcomes between the assessed therapies.

Although no official threshold is recognized by health authorities, the resulting ICUR was compared with the latest willingness-to-pay threshold estimated for the Spanish setting (€20,000/QALY) [39].

| Health state                          | Average utility (SD) [26] | HR mortality vs. general population (95% CI) [14] |
|---------------------------------------|---------------------------|---------------------------------------------------|
| AF (basal)                            | 0.727 (0.0095)            | 1.34 (1.20–1.53)                                  |
| Mild stroke                           | 0.6151 (0.0299)           | 3.18 (1.82–4.92)                                  |
| Moderate stroke                       | 0.5646 (0.0299)           | 5.84 (4.08–7.60)                                  |
| Severe stroke                         | 0.5142 (0.0299)           | 15.75 (13.99–17.51)                               |
| Systemic embolism                     | 0.6265 (0.0299)           | 1.34 (1.20–1.53)                                  |
| Myocardial infarction, males          | 0.6098 (0.0299)           | 2.56 (3.44–5.03)                                  |
| Myocardial infarction, females        |                          | 4.16 (2.27–2.88)                                  |
| Utility decrements related to complications | Utility decrement [26]   | Decrement duration                               |
| Intracranial haemorrhage              | 0.1511                    | (6 weeks)                                         |
| Other major bleedings                 | 0.1511                    | (2 weeks)                                         |
| Clinically relevant non-major bleedings | 0.0582                  | (2 days)                                          |
| Other cardiovascular hospital admissions | 0.1276                  | (6 days)                                          |
| Utility decrements associated with the use of anticoagulation (apixaban or edoxaban) | 0.0020 | 

AF atrial fibrillation, CI confidence interval, HR hazard ratio, SD standard deviation

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Sensitivity Analyses

Probabilistic sensitivity analysis (SA), to account for variability in outcomes due to statistical uncertainty in inputs, as well as one-way SA, to examine the effects of changes in key model parameters, was also performed.

A set of one-way SAs was performed by varying the base-case values of the parameters with uncertainty. The one-way SA comprised modifications of the discount rate (0%; 5%), mean patient age according to the interquartile range of patients in the ARISTOTLE trial [8] (63–77 years), economic inputs such as event management costs and resource costs (± 10% and ± 20%) and values concerning clinical inputs (event risks, HRs, death and case fatality rates and utilities) according to the 95% confidence intervals and standard deviations, if available, or assuming a standard error of 25% of the mean value.

The probabilistic SA consisted of simultaneous variation of all the potentially relevant parameters according to a distribution function previously assigned and adjusted to the data variability. The variations were conducted by running 2000 Monte Carlo iterations over the assessed 1000-patient cohort entering the model. The parametric functions applied were the beta distribution for utilities, the gamma

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| Table 3  Unitary costsa |
|--------------|----------------|----------------|
| Costs        | Retail price plus VAT [28] | Daily cost (retail price plus VAT) |
| Pharmaceutical cost |                      |                |
| Apixaban 5 mg 28 tablets | 42.40 | 3.03 |
| Edoxaban 30 mg 28 tablets | 81.30 | 2.90 |
| Edoxaban 60 mg 28 tablets | 81.30 | 2.90 |
| Event costsb | Acute phase (per episode) | Maintenance (per month) |
| Mild stroke (excluding haemorrhagic stroke). Event cost (DRG 14: stroke with infarction) + 2-week mild stroke maintenance cost | 5656.42 (4331–7156) [34, 37] | 132.76 (102–168) [34] |
| Moderate stroke (excluding haemorrhagic stroke). Event cost (DRG 14: stroke with infarction) + 2-week moderate stroke maintenance cost | 6000.82 (4595–7591) [34, 37] | 821.56 (629–1039) [34] |
| Severe stroke (excluding haemorrhagic stroke). Event cost (DRG 14: stroke with infarction) + 2-week severe stroke maintenance cost | 6741.66 (5162–8528) [34, 37] | 2303.23 (1764–2914) [34] |
| Fatal ischaemic stroke (DRG 14: Stroke with infarction) | 5590.04 (per episode) (4281–7072) [37] |                |
| Mild haemorrhagic stroke. Event cost (DRG 810: ICH) + 2-week mild stroke maintenance cost | 8024.64 (6145–10,152) [34, 37] | 132.76 (102–168) [34] |
| Moderate haemorrhagic stroke. Event cost (DRG 810: ICH) + 2-week moderate stroke maintenance cost | 8348.07 (6392–10,561) [34, 37] | 821.56 (629–1039) [34] |
| Severe haemorrhagic stroke. Event cost (DRG 810: ICH) + 2-week severe stroke maintenance cost | 9043.82 (6925–11,441) [34, 37] | 2303.23 (1764–2914) [34] |
| Fatal haemorrhagic stroke (DRG 810: ICH) | 7962.29 (per episode) (6097–10,073) [37] |                |
| Systemic embolism (DRG 131: peripheral vascular disorder without complications) | 2798.44 (2143–3540) [37] | 121.34 (93–153) [31] |
| Other ICH (excluding haemorrhagic stroke) (DRG 810: ICH) | 7962.29 (per episode) (6097–10,073) [37] |                |
| Other major bleedings (excluding ICH) | 3796.84 (per episode) (2907–4803) [35] |                |
| Clinically relevant non-major bleedings | 2549.72 (per episode) (1952–3225) [35] |                |
| MI (DRG 123: Acute MI) | 9699.48 (7427–12,270) [37] | 171.72 (131–217), estimated from [32, 33] |
| Other CV hospitalization (DRG 15: Non-specific cerebrovascular disorder) | 3800.21 (per episode) (2910–4807) [37] |                |
| Other costs | Acute phase (per episode) | Maintenance (per month) |
| Routine care | 37.88 per visit (29–48) [37] |                |
| Renal monitoring | 3.02 (NA) [37] |                |
| Dyspepsia management | 28.91 (NA) [36] |                |

CV cardiovascular, DRG diagnosis-related group, ICH intracranial haemorrhage, MI myocardial infarction, NA not applicable, VAT value-added tax
aCosts are presented as €, year 2019 values
bCosts are presented as mean (95% confidence interval)
distribution for costs and the HR of death and the log-normal distribution for the HR of clinical events.

3 Results

At the end of the simulation, for a cohort comprising 1000 patients with AF, apixaban 5 mg bid could additionally avoid five ischaemic strokes, one other intracranial haemorrhage, two MIs, six major bleedings and 29 clinically relevant non-major bleedings compared with those estimated for a 1000-patient cohort treated with edoxaban 60 mg daily. No differences were observed in the number of haemorrhagic strokes, recurrent haemorrhagic strokes or SEs. One additional recurrent ischaemic stroke in the 1000-patient cohort was reported for apixaban versus edoxaban 60 mg.

In comparison with 1000 patients treated with edoxaban 30 mg daily, the number of additional avoided events with apixaban 5 mg bid would be 21 ischaemic strokes, two SEs and eight MIs. There were no differences in terms of recurrent haemorrhagic strokes or other intracranial haemorrhages. In comparison with edoxaban 30 mg, apixaban was associated with more haemorrhagic strokes (seven additional cases) and other bleedings (48 additional major bleedings and 17 additional clinically relevant non-major bleedings) in the 1000-patient cohort. The complete detailed results are shown in Table 4.

In terms of quality-adjusted survival, apixaban would yield 6.924 QALYs per patient during the lifetime horizon compared with 6.882 QALYs yielded by edoxaban 60 mg and 6.844 QALYs by edoxaban 30 mg.

The estimated total cost per patient for each of the therapeutic alternatives from the NHS perspective would be €19,053 for apixaban, €18,651 for edoxaban 60 mg and €19,025 for edoxaban 30 mg.

From a societal perspective, the total cost per patient would be €32,620 for apixaban 5 mg bid compared with €32,297 and €34,034 for edoxaban 60 mg and 30 mg, respectively.

The ICUR from the NHS perspective was €9639 per QALY gained with apixaban versus edoxaban 60 mg and €354 per QALY gained with apixaban versus edoxaban 30 mg. From the societal perspective, the ICUR was €7757 per additional QALY with apixaban 5 mg bid versus edoxaban 60 mg daily, whereas apixaban 5 mg bid was a dominant alternative (more effective, less costly) versus edoxaban 30 mg daily.

Figure 2 is a graphical representation of the deterministic SA performed, including results for one-way SA over the discount rate, mean patient age, event and resource costs, and the ten clinical parameters that showed the greatest variations in ICUR versus the base-case results from the NHS perspective.

For apixaban compared with edoxaban 60 mg, the greatest variation was found for the stroke HR for edoxaban 60 mg, followed by the HR for the mortality trial period for edoxaban 60 mg. ICURs for the rest of the SAs were below €25,000/QALY. For apixaban compared with edoxaban 30 mg, stroke HR was also the parameter that had the widest variation, but the ICUR was lower than €15,000/QALY.

Probabilistic SA results are plotted in the cost-effectiveness planes in Fig. 3a for apixaban versus edoxaban 60 mg and Fig. 3b for apixaban compared with edoxaban 30 mg. In both comparisons, most of the iterations fall in the south-eastern quadrant, although a certain number of simulations appeared in the western quadrants, indicating cases when apixaban was a less effective option than edoxaban.

Higher dispersion of the iterations was observed in the probabilistic SA for apixaban versus edoxaban 60 mg than for apixaban versus edoxaban 30 mg. The lower treatment discontinuation rates applied for edoxaban 30 mg (HR vs. apixaban 1.04; 95% confidence interval (CI) 0.96–1.13) than for edoxaban 60 mg (HR vs. apixaban 1.10; 95% CI 1.01–1.19) led to a decreased probability of apixaban being associated with negative incremental costs.

From the NHS perspective, 71% and 74% of the iterations of apixaban versus edoxaban 60 mg in the probabilistic SA yielded an ICUR below a willingness-to-pay threshold of €20,000/QALY and €30,000/QALY, respectively (Fig. 4a).

Apixaban compared with edoxaban 30 mg was a cost-effective option in 86% and 88% of the iterations, considering willingness-to-pay thresholds of €20,000 and €30,000 per QALY gained, respectively (Fig. 4b).

4 Discussion

The present analysis complements a series of cost-effectiveness analyses performed for apixaban compared with other oral anticoagulant therapies for the prevention of stroke and SE in Spanish patients with AF. All of these economic evaluations were based on models with characteristics similar to those used in international evaluations.

In our analysis, apixaban was a cost-effective option in comparison with edoxaban 60 mg daily (€9639/QALY) and versus edoxaban 30 mg daily (€354/QALY) from the NHS perspective. An equivalent price was established for both dosages of edoxaban. Since edoxaban 30 mg was associated with a higher HR for stroke, MI and SE than edoxaban 60 mg, a lower ICUR was found for apixaban versus edoxaban 30 mg than for apixaban versus edoxaban 60 mg. Since this is the only economic evaluation assessing apixaban versus edoxaban in Spain, no comparison with other local results was possible. Our results are consistent with those obtained in previously published...
publications, which concluded that apixaban was cost effective versus dabigatran 110 mg [16], acenocoumarol [15], rivaroxaban [17] and ASA [25]. All ICURs estimated in these studies were also below the latest willingness-to-pay threshold estimated for the Spanish setting [39].

The economic evaluation of apixaban versus edoxaban in the UK [14] provided conclusions in the same kind of evaluation that was performed for the Spanish setting. Apixaban was a dominant option in comparison with low-dose edoxaban (30 mg) and a cost-effective alternative versus edoxaban 60 mg, with an ICUR of £6703/QALY gained (year 2012 values)

The present analysis is not exempt from limitations. The theoretical nature, inherent in every model, may not exactly represent usual clinical practice. The validity of analytic decision models depends on the quality of the data they are based on. In this case, the efficacy source included in the model was an indirect-comparison meta-analysis because of the lack of direct comparisons between the alternatives evaluated. This methodology, when developed with

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Table 4  Base-case results

| Base-case results                        | API       | EDO 30 mg | Diff. API vs. EDO 30 mg | EDO 60 mg | Diff. API vs. EDO 60 mg |
|-----------------------------------------|-----------|-----------|-------------------------|-----------|-------------------------|
| No. of events (in total population)     |           |           |                         |           |                         |
| Ischaemic stroke                        | 248       | 269       | −21                     | 253       | −5                      |
| Recurrent ischaemic stroke              | 22        | 24        | −2                      | 21        | 1                       |
| Haemorrhagic stroke                     | 28        | 21        | −7                      | 28        | −                       |
| Recurrent haemorrhagic stroke           | 2         | 2         | −                       | 2         | −                       |
| Systemic embolism                       | 26        | 28        | −2                      | 26        | −                       |
| Other ICH                               | 13        | 13        | 0                       | 14        | −1                      |
| Other major bleedings                   | 176       | 127       | 49                      | 182       | −6                      |
| CRNM bleeding                           | 308       | 291       | 17                      | 337       | −29                     |
| Myocardial infarction                   | 91        | 99        | −8                      | 93        | −2                      |
| Other cardiovascular hospitalization    | 1270      | 1237      | 33                      | 1267      | 3                       |
| Death                                   | 998       | 998       | −                       | 998       | −                       |
| Outcomes (per patient)                  |           |           |                         |           |                         |
| LYG                                     | 9.767     | 9.678     | 0.089                   | 9.711     | 0.056                   |
| QALYs                                   | 6.924     | 6.844     | 0.080                   | 6.882     | 0.042                   |
| Costs (per patient)                     |           |           |                         |           |                         |
| Total costs (NHS)                       | 19,053.35 | 19,025.13 | 28.23                   | 18,651.40 | 401.95                  |
| Anticoagulant drug acquisition          | 4890.50   | 4614.68   | 275.82                  | 4407.33   | 483.71                  |
| Events management                       | 12,433.82 | 12,695.98 | −262.16                 | 12,510.28 | −76.46                  |
| Dyspepsia management and renal monitoring | 4.39   | 4.33       | 0.06                    | 4.34       | 0.05                    |
| Clinical follow-up                      | 1724.64   | 1710.14   | 14.5                    | 1729.45   | −4.81                   |
| Total costs (societal perspective)      | 32,620.20 | 34,033.98 | −1413.78                | 32,296.75 | 323.45                  |
| Anticoagulant drug acquisition          | 4890.50   | 4614.68   | 275.82                  | 4407.33   | 483.71                  |
| Events management                       | 26,000.66 | 27,704.83 | −1704.17                | 26,155.63 | −154.97                 |
| Dyspepsia management and renal monitoring | 4.39   | 4.33       | 0.06                    | 4.34       | 0.05                    |
| Clinical follow-up                      | 1724.64   | 1710.14   | −1413.78                | 1729.45   | −4.81                   |
| ICURs and ICERs                         | API vs. EDO 30 mg | API vs. EDO 60 mg |
| ICURb, NHS perspective                  | 354.22    | 9639.33   |                         |           |                         |
| ICURb, societal perspective             | Dominant  | 7756.62   |                         |           |                         |
| ICERc, NHS perspective                  | 318.88    | 7261.15   |                         |           |                         |
| ICERc, societal perspective             | Dominant  | 5842.94   |                         |           |                         |

API apixaban, CRNM clinically relevant non-major, Diff. difference, EDO edoxaban, ICER incremental cost-effectiveness ratio, ICH intracranial haemorrhage, ICUR incremental cost/utility ratio, LYG life-year gained, NHS national health service, QALY quality-adjusted life-year

aCosts are presented as €, year 2019 values
bCost per QALY
cCost per LYG
transparency and rigour, is widely recognized and accepted by health technology assessment agencies, such as the UK National Institute for Health and Care Excellence [40], and other scientific associations, such as the GENESIS group, belonging to the Spanish Society of Hospital Pharmacists [41]. The pivotal trials on which authorizations were based (ARISTOTLE [8] and ENGAGE-AF [12]) were considered the most robust studies to use as the source of individual efficacies. Additionally, the common comparator used in the trials (warfarin) allowed us to perform an indirect comparison by the Bucher method.

This indirect-comparison meta-analysis did not consider the potential influence of the patients’ baseline characteristics. Although patient age is a potential key driver, the results of the one-way SA indicated that age was not among the factors that had a major influence on the cost-effectiveness results.

The assumption of allowing one complication in each cycle is not representative of clinical practice, but it was necessary given the data availability. In the same sense, facing the absence of evidence for edoxaban, an equivalent frequency of dyspepsia was applied to both alternatives; again, the SA indicated that this assumption did not influence the results.

European Society of Cardiology guidelines [5] no longer recommend ASA monotherapy in patients with AF for stroke prevention, noting that it actually causes harm, incurs a higher bleeding risk and has very little effect on preventing thromboembolic events; however, its administration as a second-line treatment was implemented in the model for patients who stopped treatment. Given the lower cost of ASA, no significant influence on the results was anticipated. However, this assumption allowed us to model patients’ transitions, at least with the ASA efficacy reported following the SA description is below the ICUR estimated in the BC. The solid grey area in bars indicates that the ICUR of the SA with the second value specified in the parentheses following the SA description is above the ICUR estimated in the BC. ASA acetylsalicylic acid, BC base case, CRNMB clinically relevant non-major bleeding, CV cardiovascular, HR hazard ratio, ICH intracranial haemorrhage, ICUR incremental cost/utility ratio, QALY quality-adjusted life-year, SA sensitivity analysis. *Apixaban was a less effective option than edoxaban 60 mg.
in the AVERROES trial [9], faced with the lack of published evidence on the use of other direct-acting oral anticoagulants as second-line treatments.

Utility values were obtained from published literature given that the apixaban clinical studies for this indication did not include quality-of-life assessment questionnaires that allowed the collection of appropriate utility values for use in this economic evaluation. The values used in this analysis were obtained from patients in the UK because no specific data for the Spanish population were available. Utility values were endorsed by an expert panel that considered these data to be representative of Spanish patients. This supports the idea that utility values derived from health states (by using defined values from the EQ-5D questionnaire) do not differ between the general populations of different European countries [42].

The societal perspective included both medical and non-medical costs but not indirect costs because of the
lack of reliable data about productivity loss in the Spanish AF population. This issue was not expected to have an impact on the results because of the advanced average age of the patients (70 years) in the simulated cohort.

Despite the limitations, both deterministic and probabilistic SAs confirmed the robustness of the model, as the vast majority of the estimated resulting ICURs did not show great variations with respect to the ICURs of the base cases.

5 Conclusion

This is the first economic evaluation to compare apixaban versus edoxaban in the Spanish setting. The methodology complied with international recommendations for the development of economic evaluations, and the results were concordant with previously published figures. Being the latest in a series of cost-effectiveness analyses for
apixaban, it provides interesting information to be considered during the medical decision-making process. This study showed that apixaban 5 mg bid is a cost-effective option for the treatment of patients with AF in Spain compared with edoxaban 30 or 60 mg daily.

Author Contributions  CS, JLLS, JRGJ validated the parameters and provided data about local management of patients. IO and FAN performed the analysis. All the authors participated in the interpretation of the results. All authors had complete access to the data. IO and FAN drafted the manuscript. All the authors reviewed and approved the final manuscript.

Data Availability  The datasets generated and/or analysed during the current study are available from the corresponding author on reasonable request.

Compliance with Ethical Standards

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Conflict of interest  IO and FAN are employed at Pharmacoeconomics and Outcomes Research Iberia (PORIB), a consultant company that specializes in economic evaluation of health technologies and has received financial support from Bristol-Myers Squibb S.A.U. for the development of the present work. CS, JLLS and JRGJ have received an unrestricted grant, as members of an advisory board, to validate the model inputs. JSuarez and CP are full-time employees of Bristol-Myers Squibb S.A.U. JSoto is a full-time employee of Pfizer S.L.U.

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