# 1 Original Article

# <sup>2</sup> Edoxaban versus Vitamin K Antagonist

- <sup>3</sup> for Atrial Fibrillation after TAVR
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# 1 Abstract

### 2 Background

3 The role of direct oral anticoagulants as compared with vitamin K antagonists

- 4 for atrial fibrillation after successful transcatheter aortic-valve replacement
- 5 (TAVR) remains unclear.

# 6 Methods

7 We conducted a multicenter, prospective, randomized, open-label, adjudicator-8 masked trial comparing edoxaban with vitamin K antagonists in patients with prevalent or incident atrial fibrillation as the indication for oral anticoagulation 9 after successful TAVR. The primary efficacy outcome was a composite of adverse 10 events consisting of death from any cause, myocardial infarction, ischemic 11 stroke, systemic thromboembolism, valve thrombosis, or major bleeding. The 12 primary safety outcome was major bleeding. On the basis of a hierarchical 13 testing plan, the primary efficacy and safety outcomes were tested sequentially 14 for noninferiority, with noninferiority of edoxaban established if the upper 15 boundary of the 95% confidence interval for the hazard ratio did not exceed 16 1.38. Superiority testing of edoxaban for efficacy would follow if noninferiority 17 and superiority were established for major bleeding. 18

### 19 Results

20 A total of 1426 patients were enrolled (713 in each group). The mean age of

21 the patients was 82.1 years, and 47.5% of the patients were women. Almost

22 all the patients had atrial fibrillation before TAVR. The rate of the primary

23 efficacy outcome was 17.3 per 100 person-years in the edoxaban group and 16.5

24 per 100 person-years in the vitamin K antagonist group (hazard ratio, 1.05;

25 95% confidence interval [CI], 0.85 to 1.31; P=0.01 for noninferiority). Rates

26 of major bleeding were 9.7 per 100 person-years and 7.0 per 100 person-years,

27 respectively (hazard ratio, 1.40; 95% CI, 1.03 to 1.91; P=0.93 for noninferiority);

28 the difference between groups was mainly due to more gastrointestinal bleeding

29 with edoxaban. Rates of death from any cause or stroke were 10.0 per 100

30 person-years in the edoxaban group and 11.7 per 100 person-years in the

31 vitamin K antagonist group (hazard ratio, 0.85; 95% CI, 0.66 to 1.11).

# 32 Conclusions

In patients with incident or prevalent atrial fibrillation who had successfulTAVR, edoxaban was noninferior to vitamin K antagonists as determined by

35 a hazard ratio margin of 38% for a composite primary outcome of adverse

36 clinical events. The incidence of major bleeding was higher with edoxaban than

37 with vitamin K antagonists. (Funded by Daiichi Sankyo; ENVISAGE-TAVI AF

38 ClinicalTrials.gov number, NCT02943785.)

39 Atrial fibrillation occurs in approximately 33% of patients after transcatheter

- 40 aortic-valve replacement (TAVR),<sup>1-7</sup> and oral anticoagulation is generally
- 41 recommended as treatment. Non-vitamin K oral anticoagulants are frequently
- 42 used for this purpose instead of vitamin K antagonists.<sup>8</sup> The effects of various

1 antithrombotic strategies to prevent thromboembolic events with atrial

2 fibrillation after TAVR have not been well studied. A randomized trial9 showed

3 that the addition of clopidogrel to oral anticoagulation in patients undergoing

4 TAVR who had established indications for anticoagulation, predominantly

5 atrial fibrillation, resulted in more bleeding complications. Non-vitamin K oral

6 anticoagulants were prescribed in less than 33% of the patients in that trial;

7 there was no comparison between regimens, and medications were mostly

### 8 initiated before TAVR.9

Edoxaban is an oral, reversible, direct factor Xa inhibitor that was shown to 9 be noninferior to a vitamin K antagonist (warfarin) in the prevention of stroke 10 and other thromboembolic events, with lower rates of bleeding and death from 11 cardiovascular causes, in a general population of patients with atrial fibrillation 12 who were at moderate-to-high thromboembolic risk, but that trial did not 13 include patients with TAVR.<sup>10</sup> An exploratory subgroup analysis involving 191 14 patients with previous implantation of a bioprosthetic valve, the results of which 15 are reported in a separate article,<sup>11</sup> suggested that clinical outcomes may have 16 been better with edoxaban than with warfarin. The aim of the current Edoxaban 17 versus Standard of Care and Their Effects on Clinical Outcomes in Patients 18 Having Undergone Transcatheter Aortic Valve Implantation-Atrial Fibrillation 19 (ENVISAGE-TAVI AF) trial was to compare the efficacy and safety of edoxaban 20 with those of vitamin K antagonists in patients with prevalent or incident atrial 21

22 fibrillation after successful TAVR.

# 23 Methods

## 24 Trial Design and Oversight

25 This trial was a multinational, multicenter, prospective, randomized, open26 label, adjudicator-masked trial.<sup>12</sup> It was conducted in accordance with the
27 International Council for Harmonisation and the Declaration of Helsinki. The
28 protocol (available with the full text of this article at NEJM.org) was approved by
29 the ethics committees and corresponding health authorities for all sites. All the
30 patients provided written informed consent before enrollment.

The sponsor, Daiichi Sankyo, contributed to the trial design, conduct, and 31 oversight; data analysis; and manuscript writing. The trial was designed by eight 32 academic authors and one author employed by the sponsor. The manuscript was 33 written by the first three authors and the last author. Confidentiality agreements 34 were in place between all the authors and the sponsor. Academic authors were 35 not restricted in publishing the data. The sponsor covered all costs associated 36 with the trial, including the cost of the anticoagulants and all tests for trial 37 purposes that were not otherwise clinically indicated. Most data analyses were 38 performed by a clinical research organization (Covance) and paid for by the 39 sponsor. All events were documented from sources, including, but not limited 40 to, paper and electronic charts, laboratory and imaging test reports, and death 41 certificates, and were adjudicated by an independent clinical events committee, 42 43 whose members were unaware of the trial-group assignments. Serious adverse

1 events were reviewed by an independent data and safety monitoring board

2 according to a predefined schedule.

# 3 Patient Selection and Randomization

4 Patients 18 years of age or older with either prevalent or incident atrial

5 fibrillation lasting more than 30 seconds after successful TAVR for severe aortic

<sup>6</sup> stenosis were eligible for enrollment. Successful TAVR was defined as correct

7 positioning of any approved transcatheter bioprosthetic aortic valve into the

8 proper anatomical location with the intended valve performance and without

9 unresolved periprocedural complications. Among the key exclusion criteria

10 were coexisting conditions that confer a high risk of bleeding (Table S1 in the

11 Supplementary Appendix, available at NEJM.org).

12 Randomization was stratified according to the use or nonuse{q1} of a

13 coronary stent for which the patient required antiplatelet medication and was

14 performed by means of an interactive Web-response system. Patients were

15 randomly assigned in a 1:1 ratio to receive edoxaban or a vitamin K antagonist

16 (any of the following drugs according to country availability: warfarin,

17 phenprocoumon, acenocoumarol, or fluindione). Randomization occurred 12

18 hours to 7 days after TAVR.

## 19 Trial Treatment and Follow-up

20 The edoxaban group received 60 mg once daily; a creatinine clearance

21 (Cockcroft–Gault formula) of 15 to 50 ml per minute, a body weight of 60 kg

22 or less, and the use of certain P-glycoprotein inhibitors were indications for

23 dose adjustment to 30 mg once daily. Edoxaban was supplied by the sponsor to

24 the sites, and vitamin K antagonists were supplied according to local practice.

25 The target international normalized ratio (INR) for the vitamin K antagonist

26 regimen was 2.0 to 3.0 (adjusted to 1.6 to 2.6 for patients ≥70 years of age in

27 Japan). Specified antiplatelet therapy in either trial group (stratification variable)

<sup>28</sup> was allowed at the treating physician's discretion, including dual antiplatelet

29 therapy for up to 3 months or single antiplatelet therapy indefinitely. Patients

30 were followed at 3 months after randomization and every 6 months thereafter

31 (minimum of 6 months up to 36 months); details regarding follow-up and

32 concomitant medications are provided in Section 3 in the Supplementary

33 Appendix and in the protocol.

## 34 Outcomes

35 The primary efficacy outcome was the incidence of net adverse clinical events,

<sup>36</sup> defined as the composite of death from any cause, myocardial infarction,

37 ischemic stroke, systemic thromboembolic event, valve thrombosis, or major

38 bleeding (International Society on Thrombosis and Haemostasis [ISTH]

39 definition).<sup>13</sup> The primary safety outcome was the incidence of major bleeding,

40 designated according to ISTH definitions as clinically overt bleeding associated

41 with a reduced hemoglobin level, blood transfusion, symptomatic bleeding

<sup>42</sup> at a critical site, or death.<sup>13</sup> Secondary outcomes were bleeding as defined by the Global Utilization of Streptokinase and Tissue Plasminogen Activator for 1 Occluded Coronary Arteries (GUSTO), Thrombolysis in Myocardial Infarction

2 (TIMI), and Bleeding Academic Research Consortium (BARC), as well as

3 components of the composite primary outcome; secondary efficacy and safety

4 outcomes are listed in Sections 4.1.2 and 4.2.2 in the statistical analysis plan,

5 available with the protocol. Clinically relevant nonmajor bleeding was defined

6 according to ISTH criteria.<sup>13</sup> (Details on outcome and bleeding definitions are

7 provided in Tables S2 and S3.)

## 8 Statistical Analysis

We estimated that 320 events would be needed in approximately 1400 patients 9 to show noninferiority of edoxaban to vitamin K antagonists for the primary 10 outcome with 80% power and a two-sided significance level of 0.05; details on 11 power calculation and statistics are provided in the statistical analysis plan. 12 The primary analyses were performed in the intention-to-treat population. A 13 four-step hierarchical testing strategy was used, sequentially testing edoxaban 14 as compared with vitamin K antagonists for noninferiority for the primary 15 outcome, noninferiority for major bleeding, superiority for major bleeding, and 16 superiority for the primary outcome. For both the primary efficacy and major 17 bleeding outcomes, noninferiority would be established if the upper boundary 18 of the 95% confidence interval for the hazard ratio did not exceed 1.38 (Fig. 19 S1). Superiority testing was based on a two-sided significance level of 0.05. The 20 primary analysis period was the time from randomization to an end-of-treatment 21

21 primary analysis period was the time from randomization to an end-of-treatment

22 visit at 36 months, an end-of-trial visit, the patient's last visit, or death,

23 whichever occurred first.

Cumulative event-free survival was estimated by means of Kaplan-Meier 24 analyses. Cox proportional-hazards regression models were used to analyze 25 the time from randomization to the first occurrence of a trial outcome, with 26 treatment regimen as a main factor and two randomization stratification 27 factors (coronary stent for which the patient required antiplatelet medication 28 and characteristics warranting adjustment of the edoxaban dose) as covariates, 29 to estimate the hazard ratios and 95% confidence intervals. The proportionality 30 assumption was tested by visual inspection of the log-minus-log survival curves 31 of the outcomes, and the proportionality assumption was upheld. 32 Secondary outcomes were analyzed with the use of the same methods as 33 those described above, with comparisons focused on superiority but without 34 hierarchical analysis. Because of the lack of a prespecified plan for adjustment 35 of confidence intervals for multiple comparisons, no conclusions can be drawn 36 from the secondary outcome results. All other safety outcomes were summarized 37 with the use of descriptive statistics (SAS software, version 9.2 or newer; SAS 38 Institute). Detailed descriptions of all statistical analyses are provided in the 39 statistical analysis plan. 40

# 1 Results

# 2 Trial Population

3 From April 2017 through January 2020, a total of 1426 patients with prevalent or

4 incident atrial fibrillation and conventional indications for oral anticoagulants
5 were enrolled after successful TAVR at 173 centers in 14 countries on three

6 continents (Fig. 1 and Table S4). A total of 713 patients were assigned to

7 the edoxaban group, and an equal number were assigned to the vitamin K

8 antagonist group. Almost all the patients had atrial fibrillation before TAVR. The

9 mean time between TAVR and randomization was 66.6 hours in the edoxaban

10 group and 70.2 hours in the vitamin K antagonist group. The demographic and

11 clinical characteristics of the patients at baseline were similar in the two trial

12 groups (Table 1). The mean age of the patients was 82.1 years, and 47.5% of

13 the patients were women. The mean Society of Thoracic Surgeons risk score

14 (predicted 30-day mortality) was 4.9%, and the mean CHA<sub>2</sub>DS<sub>2</sub>-VASc score

15 (range, 0 to 9, with higher scores indicating greater risk of embolic events) was

16 4.5. There was concomitant use of oral antiplatelet agents before randomization

17 in 328 patients (46.0%) in the edoxaban group and in 359 patients (50.4%) in the

18 vitamin K antagonist group. At trial entry, 46.4% of the overall trial population

met any of the criteria for adjustment of the edoxaban dose and received reduceddoses.

The median duration of follow-up was 554 days in the edoxaban group and

22 530 days in the vitamin K antagonist group. The mean and median percent of

23 time of INR within the therapeutic range in the vitamin K antagonist group

24 were 63.5% and 68.2%, respectively (Fig. S2). During the entire trial period, 215

25 patients (30.2%) in the edoxaban group discontinued the trial drug, as compared

26 with 289 patients (40.5%) in the vitamin K antagonist group (Table S5 and Fig.

27 S3). Use of concomitant antiplatelet therapy is summarized in Table S6.

## 28 Efficacy and Safety Outcomes

In the intention-to-treat analysis, a net adverse clinical event (primary efficacy 29 outcome) occurred in 170 patients (17.3 per 100 person-years) in the edoxaban 30 group and in 157 patients (16.5 per 100 person-years) in the vitamin K 31 antagonist group (hazard ratio, 1.05; 95% confidence interval [CI], 0.85 to 1.31; 32 noninferiority margin, 1.38; P=0.01 for inferiority) (Table 2 and Figs. 2 and 3). 33 Major bleeding (primary safety outcome) occurred in 98 patients (9.7 per 100 34 person-years) in the edoxaban group and in 68 patients (7.0 per 100 person-35 years) in the vitamin K antagonist group (hazard ratio, 1.40; 95% CI, 1.03 to 36 1.91; noninferiority margin, 1.38; P=0.93 for noninferiority). The hierarchical 37 testing failed at this step; hence, formal testing for superiority was not 38 performed. 39

The rate of intracranial hemorrhage was 1.5 per 100 person-years in the edoxaban group and 2.1 per 100 person-years in the vitamin K antagonist group, and the rate of fatal bleeding was 1.0 per 100 person-years in both trial groups (Table 2). More patients in the edoxaban group than in the vitamin K antagonist 1 group had major gastrointestinal bleeding (56 [5.4 per 100 person-years] vs.

2 27 [2.7 per 100 person-years]; hazard ratio, 2.03; 95% CI, 1.28 to 3.22), despite

3 similar incidences of administration of proton-pump inhibitors (71.7% and

4 69.0%, respectively); one case of major gastrointestinal bleeding was fatal in the

5 edoxaban group. Among patients with major gastrointestinal bleeding, 46 of 56

6 (82%) in the edoxaban group and 26 of 27 (96%) in the vitamin K antagonist

7 group received proton-pump inhibitors.

Death from any cause occurred in 85 patients (7.8 per 100 person-years) 8 in the edoxaban group and in 93 patients (9.1 per 100 person-years) in the 9 vitamin K antagonist group (hazard ratio, 0.86; 95% CI, 0.64 to 1.15). The 10 rate of ischemic stroke was 2.1 per 100 person-years and 2.8 per 100 person-11 years, respectively (hazard ratio, 0.75; 95% CI, 0.43 to 1.30). The rate of 12 myocardial infarction was 1.1 per 100 person-years and 0.7 per 100 person-years, 13 respectively. Systemic thromboembolic events were rare (0.2 per 100 person-years 14 in the edoxaban group and 0.3 per 100 person-years in the vitamin K antagonist 15 group), and no cases of valve thrombosis occurred (Table 2). The rate of death 16 from any cause or stroke was 10.0 per 100 person-years in the edoxaban group 17 and 11.7 per 100 person-years in the vitamin K antagonist group (hazard ratio, 18 0.85; 95% CI, 0.66 to 1.11). Composite outcome measures, including major 19 adverse cardiac and cerebrovascular events, are reported in Table 2. 20 The results of prespecified subgroup analyses are provided in Figures S4 21

and S5; however, the trial was underpowered for these analyses, and results
are exploratory. Secondary efficacy and safety results are shown in Tables S7
and S10, respectively. The results regarding the two primary outcomes were
concordant when other bleeding scales were applied; anticoagulation after trial
treatment is summarized in Table S8. Causes of death are reported in Table S9.

# 27 Concomitant Antiplatelet Drugs and Edoxaban Dose Adjustment

A prespecified exploratory analysis compared treatment effects of edoxaban 28 with those of vitamin K antagonists in patients with or without criteria for 29 adjustment of the edoxaban dose. Rates of net adverse clinical events were 30 similar in the edoxaban group and the vitamin K antagonist group, regardless 31 of whether these criteria were met (Fig. S6). Among patients who met these 32 criteria, rates of major bleeding were similar in the edoxaban group and the 33 vitamin K antagonist group (9.7 per 100 person-years and 7.9 per 100 person-34 vears, respectively; hazard ratio, 1.25; 95% CI, 0.80 to 1.97). Rates of death from 35 any cause were 8.1 per 100 person-years in the edoxaban group and 12.7 per 36 100 person-years in the vitamin K antagonist group (hazard ratio, 0.64; 95% 37 CI, 0.43 to 0.96); rates of death from noncardiovascular causes were 3.3 per 100 38 person-years and 6.4 per 100 person-years, respectively (hazard ratio, 0.52; 95% 39 CI, 0.28 to 0.96). Among patients who did not meet these criteria, rates of major 40 bleeding were 9.7 per 100 person-years in the edoxaban group and 6.3 per 100 41 person-years in the vitamin K antagonist group (hazard ratio, 1.54; 95% CI, 1.00 42 to 2.35); rates of death from any cause were 7.6 per 100 person-years and 6.3 per 43 44 100 person-years, respectively (hazard ratio, 1.20; 95% CI, 0.78 to 1.85). Results

1 of exploratory 90-day landmark analyses are presented in Figs. S7 through S9.

2 Post hoc analyses of rates of the primary efficacy and safety outcomes in the

3 two trial groups according to antiplatelet therapy as prescribed at randomization

4 are shown in Fig. S10.

# 5 Discussion

The ENVISAGE-TAVI AF trial compared the efficacy and safety of edoxaban with those of vitamin K antagonists in patients with prevalent or incident 7 atrial fibrillation after successful TAVR. Edoxaban was noninferior to vitamin 8 K antagonists with respect to the composite primary efficacy outcome on the 9 basis of an upper boundary of the 95% confidence interval for the hazard ratio 10 that was less than the protocol-defined noninferiority margin of 1.38. However, 11 edoxaban failed noninferiority testing regarding the rate of major bleeding, 12 which was due mainly to more major gastrointestinal bleeding in the edoxaban 13 group. The incidences of intracranial hemorrhage or fatal bleeding were low 14 and were similar in the two trial groups. Because of the hierarchical design of 15 our statistical analysis, the failure to show noninferiority for major bleeding 16 precluded formal testing for superiority of edoxaban, but the point estimate 17 for the hazard ratio favored vitamin K antagonists and the confidence interval 18 included 1, indicating that superiority of edoxaban would not have been shown. 19 Results for composite outcome measures including major adverse cardiac and 20 cerebrovascular events were similar in the two trial groups. The patients in 21 our trial did not have valve thrombosis, which may be compatible with the 22 low incidence observed in recent trials of TAVR in low-risk patients.<sup>6,7</sup> Among 23 patients who received specified concomitant antiplatelet therapy, edoxaban 24 was associated with a higher incidence of major bleeding than vitamin K 25 antagonists. 26 The 2020 American Heart Association-American College of Cardiology 27 guideline for the management of valvular heart disease acknowledged the 28 paucity of data to support non-vitamin K oral anticoagulants for atrial 29 fibrillation within 3 months after implantation of a surgical or transcatheter 30 bioprosthetic valve.<sup>14</sup> The antithrombotic regimen after TAVR has been 31 investigated in recent randomized, controlled trials.9,15-17 A trial of intermediate-32 dose rivaroxaban in patients without an indication for oral anticoagulation but 33 who were receiving antiplatelet therapy showed increased risks of major bleeding 34 and death as compared with control.<sup>16,17</sup> A randomized trial that evaluated 35 clopidogrel in addition to oral anticoagulation after TAVR in patients with an 36 indication for oral anticoagulation showed that the combination regimen was 37 associated with more bleeding than oral anticoagulation monotherapy and had 38

39 no clinical benefits.<sup>9,18</sup> Overall, in the current trial, edoxaban was associated

40 with more cases of major bleeding than vitamin K antagonists. Subtherapeutic

41 INR values and a higher incidence of drug discontinuation in the vitamin

42 K antagonist group may have affected the bleeding outcomes. Concomitant

43 antiplatelet therapy was specified before randomization in approximately 50% of

1 the patients. A post hoc analysis showed that patients who received antiplatelet

2 therapy may have had higher bleeding rates with edoxaban than with vitamin

3 K antagonists, as opposed to similar bleeding rates among patients without

4 specified antiplatelet therapy, but these comments are exploratory only. The

5 routine use of concomitant antiplatelet therapy in addition to oral anticoagulant

<sup>6</sup> therapy is no longer recommended.<sup>19</sup> Patients who met the criteria for dose

7 adjustment during the trial and received edoxaban at a dose of 30 mg once daily

8 had similar incidences of net adverse clinical events and major bleeding as those

9 who received vitamin K antagonists.

10 Trials and studies have shown a better benefit-risk profile with non-

11 vitamin K oral anticoagulants than with vitamin K antagonists in patients

12 with nonvalvular atrial fibrillation.<sup>10,20-22</sup> However, the patient populations of

13 these trials differed from that of the ENVISAGE-TAVI AF trial with respect

14 to factors such as a younger mean age by approximately a decade, a lower

15 prevalence of heart failure, and a limited number of patients with bioprosthetic

16 valves; no direct comparisons of edoxaban and vitamin K antagonists can be

17 made between these trials and ours. Acquired von Willebrand's disease and

18 arteriovenous malformations may also contribute to gastrointestinal bleeding in

19 patients with severe aortic stenosis.<sup>23</sup>

20 Our trial had an open-label design that entailed a risk of reporting bias

21 regarding the trial outcomes. The coronavirus disease 2019 pandemic affected

22 the outpatient clinic follow-up routine and may have resulted in underassessment

23 of laboratory data and mild-to-moderate clinical events. The outcomes of death

24 and trial-drug discontinuation may have been competing risks in relation to

25 the outcomes we studied. Our trial results apply only to patients with atrial

26 fibrillation, intermediate operative risk, and symptomatic aortic stenosis, and

27 the trial involved a population of older adults who were undergoing TAVR. These

28 results may not apply to younger patients at lower operative risk, patients with

29 asymptomatic aortic stenosis, and those undergoing concomitant percutaneous

30 coronary intervention. Most patients who were enrolled in the trial had atrial

31 fibrillation before TAVR.

32 In our trial involving patients who had an indication for oral anticoagulation

33 for atrial fibrillation after successful TAVR, edoxaban was noninferior to vitamin

34 K antagonists for the composite primary outcome of adverse clinical events.

35 Edoxaban was associated with a higher risk of major bleeding than vitamin K

36 antagonists.

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the manuscript

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Figure 1. Screening, Randomization, and Treatment.

VKA denotes vitamin K antagonist.



Figure 2. Kaplan-Meier Curves for the Primary Outcomes and Other Outcomes (Intention-to-Treat Population).

Net adverse clinical events were defined as a composite of death from any cause, myocardial infarction, ischemic stroke, systemic thromboembolic event, valve thrombosis, or major bleeding (International Society on Thrombosis and Haemostasis [ISTH] definition). Insets show the same data on an enlarged y axis.



#### Figure 3. Hazard Ratio for the Primary Efficacy Outcome and Its Components (Intention-to-Treat Population).

Hazard ratios and confidence intervals for the comparison of edoxaban with VKA are based on the Cox proportional-hazards regression model including treatment group, transcatheter actic-valve replacement procedure undergone with stenting (yes or no), and indication for dose adjustment (yes or no) as covariates. Major bleeding was defined according to ISTH criteria. Two components of the composite outcome of net adverse clinical events — systemic thromboembolic event and valve thrombosis — are not shown because fewer than five patients had an event in each treatment group.

Table 1. Demographic and Clinical Characteristics of the Patients at Baseline (Intention-to-Treat Population).*				
Characteristic	Edoxaban (N=713)	Vitamin K Antagonist (N=713)		
Age — yr	82.1±5.4	82.1±5.5		
Female sex — no. (%)	347 (48.7)	331 (46.4)		
Race — no. (%)†				
Asian	92 (12.9)	89 (12.5)		
White	593 (83.2)	594 (83.3)		
Other	28 (3.9)	30 (4.2)		
Weight — kg	74.6±17.9	76.0±17.3		
Body-mass index‡	27.5±5.7	27.9±5.4		
Creatinine clearance by Cockcroft–Gault formula — ml/min	57.9±24.0	58.6±24.3		
Hypertension — no. (%)	647 (90.7)	657 (92.1)		
Diabetes mellitus — no. (%)	270 (37.9)	257 (36.0)		
Congestive heart failure — no. (%)	591 (82.9)	619 (86.8)		
NYHA class III or IV	314 (44.0)	328 (46.0)		
Mitral-valve disease — no. (%)	57 (8.0)	60 (8.4)		
History of stroke or TIA — no. (%)	123 (17.3)	116 (16.3)		
History of coronary artery disease — no. (%)	293 (41.1)	297 (41.7)		
Previous CABG	67 (9.4)	60 (8.4)		
Previous PCI	176 (24.7)	192 (26.9)		
PCI performed within 30 days before TAVR	34 (4.8)	28 (3.9)		
Previous myocardial infarction	97 (13.6)	101 (14.2)		
Incident (new onset) atrial fibrillation — no. (%)	7 (1.0)	8 (1.1)		
CHA <sub>2</sub> DS <sub>2</sub> -VASc score∫				
Mean	4.5±1.4	4.5±1.3		
Median (IQR)	4 (4–5)	4 (4–5)		
STS risk score¶				
Mean	4.8±3.5	5.0±4.1		
Distribution — %				
<4	53.0	51.5		
4–8	34.7	35.5		
>8	12.3	13.0		
Gastrointestinal disorder — no. (%)	264 (37.0)	242 (33.9)		
Previous PPI use — no. (%)	406 (56.9)	393 (55.1)		
History of labile INR — no. (%)	53 (7.4)	61 (8.6)		
Indication for dose adjustment — no. (%)**	330 (46.3)	331 (46.4)		
Valve type — no. (%)††				
Any balloon-expandable valve	342 (48.0)	335 (47.0)		
Intraannular self-expanding valve	46 (6.5)	49 (6.9)		
Supraannular self-expanding valve	325 (45.6)	328 (46.0)		

\* Plus-minus values are means ±SD. Medical history was coded with the use of the Medical Dictionary for Regulatory Activities. Percentages may not total 100 because of rounding. CABG denotes coronary-artery bypass grafting, INR international normalized ratio, IQR interquartile range, NYHA New York Heart Association, PCI percutaneous coronary intervention, PPI proton-pump inhibitor, TAVR transcatheter aortic-valve replacement, and TIA transient ischemic attack.

Race was reported by the investigator from information obtained from patient history. "Other" includes patients of another race and those who chose not to report race.

The body-mass index is the weight in kilograms divided by the square of the height in meters.

The CHA<sub>2</sub>DS<sub>2</sub>-VASc is a measure of the risk of stroke among persons with atrial fibrillation. Weighted scores are based on the presence of congestive heart failure, hypertension, diabetes mellitus, or vascular disease; a history of stroke or TIA; an age of 65 to 74 years or 75 years or older; and sex. Scores range from 0 to 9, with higher scores indicating a greater risk.

Scoring on the risk model of the Society of Thoracic Surgeons (STS) uses an algorithm that is based on the presence of coexisting illnesses in order to predict 30-day operative mortality. The STS score equals the predicted mortality expressed as a percentage.

Percentages of patients were calculated on the basis of the total number of patients with data available on the STS score.

\*\* Indications for adjustment of the edoxaban dose included a creatinine clearance of 50 ml or less per minute, a body weight of 60 kg or less (not used as an indication in U.S. patients), and concomitant therapy with a P-glycoprotein inhibitor (not used as an indication in U.S. patients).

<sup>††</sup> One patient in the vitamin K antagonist group did not report valve type.<sup>{q2</sup>}

Table 2. Efficacy and Safety Outcomes (Intention-to-Treat Population).*				
Outcome	Edoxaban (N=713)	Vitamin K Antagonist (N=713)	Hazard Ratio (95% CI)	
	no. of patients (rate per 100 person-yr)			
Primary efficacy outcome: net adverse clinical events†	170 (17.3)	157 (16.5)	1.05 (0.85–1.31)‡	
Primary safety outcome: major bleeding§	98 (9.7)	68 (7.0)	1.40 (1.03–1.91)¶	
Secondary outcomes				
Death from any cause	85 (7.8)	93 (9.1)	0.86 (0.64–1.15)	
Death from cardiovascular causes	49 (4.5)	46 (4.5)	1.00 (0.67–1.50)	
Ischemic stroke	22 (2.1)	28 (2.8)	0.75 (0.43–1.30)	
Myocardial infarction	12 (1.1)	7 (0.7)	1.65 (0.65-4.14)	
Systemic thromboembolic event	2 (0.2)	3 (0.3)	Not calculated	
Valve thrombosis∬	0	0	Not calculated	
Any stroke	29 (2.7)	35 (3.5)	0.78 (0.48–1.28)	
Major adverse cardiac or cerebrovascular event	86 (8.2)	80 (8.1)	1.02 (0.76–1.39)	
Major adverse cardiac event**	61 (5.7)	53 (5.2)	1.10 (0.76–1.58)	
Fatal bleeding§	11 (1.0)	10 (1.0)	Not calculated	
Life-threatening bleeding	17 (1.6)	19 (1.9)	Not calculated	
Intracranial hemorrhage	16 (1.5)	21 (2.1)	0.72 (0.38–1.39)	
Clinically relevant nonmajor bleeding§	164 (18.2)	142 (16.4)	1.13 (0.90–1.14)	

\* The two noninferiority tests with respect to net adverse clinical events (primary efficacy outcome) and major bleeding (primary safety outcome) were the initial two steps of hierarchical testing. Because the second step failed, no further testing on superiority was performed.

Net adverse clinical events were defined as a composite of death from any cause, myocardial infarction, ischemic stroke, systemic thromboembolic event, valve thrombosis, or major bleeding (International Society on Thrombosis and Haemostasis [ISTH] definition).

 $\div$  P=0.01 for inferiority.

The ISTH definition was used.

¶ P=0.93 for noninferiority.

Major adverse cardiac and cerebrovascular events were defined as the composite of death from cardiovascular causes (including sudden, unexplained, and unwitnessed death), myocardial infarction, stroke (ischemic, hemorrhagic, or undetermined), or repeat coronary revascularization of the target lesion.

\*\* Major adverse cardiac events were defined as the composite of death from cardiovascular causes (including sudden, unexplained, and unwitnessed death), myocardial infarction, or repeat coronary revascularization of the target lesion.

# Queries

q1. AU: according to placement or no placement?

q2. AU: Was the valve type reported by the patient or by the surgeon or investigator?

# Data Sharing Statement

Van Mieghem NM, Unverdorben M, Hengstenberg C, et al. Edoxaban versus Vitamin K Antagonist for Atrial Fibrillation after TAVR. N Engl J Med. DOI: 10.1056/NEJMoa2111016.

Question	Authors' Response
Will the data collected for your study	No
be made available to others?	
Would you like to offer context for	-
your decision?	
Which data?	_
Additional information about data	—
How or where can the data be	—
obtained?	
When will data availability begin?	—
When will data availability end?	—
Will any supporting documents be	—
available?	
Which supporting documents?	—
Additional information about	-
supporting documents	
How or where can supporting	-
documents be obtained?	
When will supporting documents	-
availability begin?	
When will supporting documents	-
availability end?	
To whom will data be available?	—
For what type of analysis or purpose?	-
By what mechanism?	
Any other restrictions?	_
Additional information	-

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