ANNEX I

SUMMARY OF PRODUCT CHARACTERISTICS
This medicinal product is subject to additional monitoring. This will allow quick identification of new safety information. Healthcare professionals are asked to report any suspected adverse reactions. See section 4.8 for how to report adverse reactions.

1. NAME OF THE MEDICINAL PRODUCT

Lixiana 15 mg film-coated tablets

2. QUALITATIVE AND QUANTITATIVE COMPOSITION

Each film-coated tablet contains 15 mg edoxaban (as tosilate).

For the full list of excipients, see section 6.1.

3. PHARMACEUTICAL FORM

Film-coated tablet.
Orange, round-shaped film-coated tablets (6.7 mm diameter) debossed with “DSC L15”.

4. CLINICAL PARTICULARS

4.1 Therapeutic indications

Prevention of stroke and systemic embolism in adult patients with nonvalvular atrial fibrillation (NVAF) with one or more risk factors, such as congestive heart failure, hypertension, age ≥ 75 years, diabetes mellitus, prior stroke or transient ischaemic attack (TIA).

Treatment of deep vein thrombosis (DVT) and pulmonary embolism (PE), and prevention of recurrent DVT and PE in adults (see section 4.4 for haemodynamically unstable PE patients).

4.2 Posology and method of administration

Posology

Prevention of stroke and systemic embolism
The recommended dose is 60 mg edoxaban once daily.
Therapy with edoxaban in NVAF patients should be continued long term.

Treatment of DVT, treatment of PE and prevention of recurrent DVT and PE (VTE)
The recommended dose is 60 mg edoxaban once daily following initial use of parenteral anticoagulant for at least 5 days (see section 5.1). Edoxaban and initial parenteral anticoagulant should not be administered simultaneously.
The duration of therapy for treatment of DVT and PE (venous thromboembolism, VTE), and prevention of recurrent VTE should be individualised after careful assessment of the treatment benefit against the risk for bleeding (see section 4.4). Short duration of therapy (at least 3 months) should be based on transient risk factors (e.g. recent surgery, trauma, immobilisation) and longer durations should be based on permanent risk factors or idiopathic DVT or PE.
For NVAF and VTE the recommended dose is 30 mg edoxaban once daily in patients with one or more of the following clinical factors:

- Moderate or severe renal impairment (creatinine clearance (CrCL) 15 - 50 mL/min)
- Low body weight ≤ 60 kg
- Concomitant use of the following P-glycoprotein (P-gp) inhibitors: ciclosporin, dronedarone, erythromycin, or ketoconazole.

**Table 1: Summary of posology in NVAF and VTE (DVT and PE)**

<table>
<thead>
<tr>
<th>Summary Guide for Dosing</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Recommended dose</strong></td>
<td>60 mg once daily</td>
</tr>
<tr>
<td><strong>Dose recommendation for patients with one or more of the following clinical factors:</strong></td>
<td></td>
</tr>
<tr>
<td>Renal Impairment</td>
<td>Moderate or severe (CrCL 15 – 50 mL/min)</td>
</tr>
<tr>
<td>Low Body Weight</td>
<td>≤ 60 kg</td>
</tr>
<tr>
<td>P-gp Inhibitors</td>
<td>Ciclosporin, dronedarone, erythromycin, ketoconazole</td>
</tr>
</tbody>
</table>

**Missed dose**

If a dose of Lixiana is missed, the dose should be taken immediately and then be continued the following day with the once-daily intake as recommended. The patient should not take double the prescribed dose on the same day to make up for a missed dose.

**Switching to and from Lixiana**

Continued anticoagulant therapy is important in patients with NVAF and VTE. There may be situations that warrant a change in anticoagulation therapy (Table 2).

**Table 2: Switching**

<table>
<thead>
<tr>
<th>Switching to Lixiana</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>From</strong></td>
<td><strong>To</strong></td>
</tr>
<tr>
<td>Vitamin K Antagonist (VKA)</td>
<td>Lixiana</td>
</tr>
<tr>
<td>Oral anticoagulants other than VKA</td>
<td>Lixiana</td>
</tr>
<tr>
<td>Parenteral anticoagulants</td>
<td>Lixiana</td>
</tr>
</tbody>
</table>
## Switching from Lixiana

<table>
<thead>
<tr>
<th>From</th>
<th>To</th>
<th>Recommendation</th>
</tr>
</thead>
</table>
| Lixiana | Vitamin K Antagonist (VKA) | There is a potential for inadequate anticoagulation during the transition from Lixiana to VKA. Continuous adequate anticoagulation should be ensured during any transition to an alternate anticoagulant.  

**Oral option:** For patients currently on a 60 mg dose, administer a Lixiana dose of 30 mg once daily together with an appropriate VKA dose.  

For patients currently on a 30 mg dose (for one or more of the following clinical factors: moderate to severe renal impairment (CrCL 15 – 50 mL/min), low body weight, or use with certain P-gp inhibitors), administer a Lixiana dose of 15 mg once daily together with an appropriate VKA dose.  

Patients should not take a loading dose of VKA in order to promptly achieve a stable INR between 2 and 3. It is recommended to take into account the maintenance dose of VKA and if the patient was previously taking a VKA or to use valid INR driven VKA treatment algorithm, in accordance with local practice.  

Once an INR ≥ 2.0 is achieved, Lixiana should be discontinued. Most patients (85%) should be able to achieve an INR ≥ 2.0 within 14 days of concomitant administration of Lixiana and VKA. After 14 days it is recommended that Lixiana is discontinued and the VKA continued to be titrated to achieve an INR between 2 and 3.  

It is recommended that during the first 14 days of concomitant therapy the INR is measured at least 3 times just prior to taking the daily dose of Lixiana to minimise the influence of Lixiana on INR measurements. Concomitant Lixiana and VKA can increase the INR post Lixiana dose by up to 46%.  

**Parenteral option:** Discontinue Lixiana and administer a parenteral anticoagulant and VKA at the time of the next scheduled Lixiana dose. Once a stable INR of ≥ 2.0 is achieved, the parenteral anticoagulant should be discontinued and the VKA continued. |
| Lixiana | Oral anticoagulants other than VKA | Discontinue Lixiana and start the non-VKA anticoagulant at the time of the next scheduled dose of Lixiana. |
Switching from Lixiana

<table>
<thead>
<tr>
<th>From</th>
<th>To</th>
<th>Recommendation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lixiana</td>
<td>Parenteral anticoagulants</td>
<td>These agents should not be administered simultaneously. Discontinue Lixiana and start the parenteral anticoagulant at the time of the next scheduled dose of Lixiana.</td>
</tr>
</tbody>
</table>

**Special populations**

**Assessment of renal function:**
- Renal function should be assessed in all patients by calculating the creatinine clearance (CrCL) prior to initiation of treatment with Lixiana to exclude patients with end stage renal disease (i.e. CrCL < 15 mL/min), to use the correct Lixiana dose in patients with CrCL 15 – 50 mL/min (30 mg once daily), in patients with CrCL > 50 mL/min (60 mg once daily) and when deciding on the use of Lixiana in patients with increased creatinine clearance (see section 4.4).
- Renal function should also be assessed when a change in renal function is suspected during treatment (e.g. hypovolaemia, dehydration, and in case of concomitant use of certain medicinal products).

The method used to estimate renal function (CrCL in mL/min) during the clinical development of Lixiana was the Cockcroft-Gault method. The formula is as follows:

- For creatinine in µmol/L:
  \[
  \frac{1.23 \times (140-\text{age [years]}) \times \text{weight [kg]} (\times 0.85 \text{ if female})}{\text{serum creatinine [µmol/L]}}
  \]
- For creatinine in mg/dL:
  \[
  \frac{(140-\text{age [years]}) \times \text{weight [kg]} (\times 0.85 \text{ if female})}{72 \times \text{serum creatinine [mg/dL]}}
  \]

This method is recommended when assessing patients’ CrCL prior to and during Lixiana treatment.

**Renal impairment**
In patients with mild renal impairment (CrCL > 50 – 80 mL/min), the recommended dose is 60 mg Lixiana once daily.

In patients with moderate or severe renal impairment (CrCL 15 – 50 mL/min), the recommended dose is 30 mg Lixiana once daily (see section 5.2).

In patients with end stage renal disease (ESRD) (CrCL < 15 mL/min) or on dialysis, the use of Lixiana is not recommended (see sections 4.4 and 5.2).

**Hepatic impairment**
Lixiana is contraindicated in patients with hepatic disease associated with coagulopathy and clinically relevant bleeding risk (see section 4.3).

In patients with severe hepatic impairment Lixiana is not recommended (see sections 4.4 and 5.2).

In patients with mild to moderate hepatic impairment the recommended dose is 60 mg Lixiana once daily (see section 5.2). Lixiana should be used with caution in patients with mild to moderate hepatic impairment (see section 4.4).
Patients with elevated liver enzymes (ALT/AST > 2 x ULN) or total bilirubin ≥ 1.5 x ULN were excluded in clinical trials. Therefore Lixiana should be used with caution in this population (see sections 4.4 and 5.2). Prior to initiating Lixiana, liver function testing should be performed.

**Body weight**
For patients with body weight ≤ 60 kg, the recommended dose is 30 mg Lixiana once daily (see section 5.2).

**Elderly**
No dose reduction is required (see section 5.2).

**Gender**
No dose reduction is required (see section 5.2).

**Concomitant use of Lixiana with P-glycoprotein (P-gp) inhibitors**
In patients concomitantly taking Lixiana and the following P-gp inhibitors: ciclosporin, dronedarone, erythromycin, or ketoconazole, the recommended dose is 30 mg Lixiana once daily (see section 4.5). No dose reduction is required for concomitant use of amiodarone, quinidine or verapamil (see section 4.5).

The use of Lixiana with other P-gp inhibitors including HIV protease inhibitors has not been studied.

**Paediatric population**
The safety and efficacy of Lixiana in children and adolescents less than 18 years of age have not been established. No data are available.

**Patients undergoing cardioversion**
Lixiana can be initiated or continued in patients who may require cardioversion. For transoesophageal echocardiogram (TEE) guided cardioversion in patients not previously treated with anticoagulants, Lixiana treatment should be started at least 2 hours before cardioversion to ensure adequate anticoagulation (see sections 5.1 and 5.2). Cardioversion should be performed no later than 12 hours after the dose of Lixiana on the day of the procedure.

**For all patients undergoing cardioversion:** Confirmation should be sought prior to cardioversion that the patient has taken Lixiana as prescribed. Decisions on initiation and duration of treatment should follow established guidelines for anticoagulant treatment in patients undergoing cardioversion.

**Method of administration**
For oral use.
Lixiana can be taken with or without food (see section 5.2).

### 4.3 Contraindications
- Hypersensitivity to the active substance or to any of the excipients listed in section 6.1.
- Clinically significant active bleeding.
- Hepatic disease associated with coagulopathy and clinically relevant bleeding risk.
- Lesion or condition, if considered to be a significant risk for major bleeding. This may include current or recent gastrointestinal ulceration, presence of malignant neoplasms at high risk of bleeding, recent brain or spinal injury, recent brain, spinal or ophthalmic surgery, recent intracranial haemorrhage, known or suspected oesophageal varices, arteriovenous malformations, vascular aneurysms or major intraspinal or intracerebral vascular abnormalities.
- Uncontrolled severe hypertension.
- Concomitant treatment with any other anticoagulants e.g. unfractionated heparin (UFH), low molecular weight heparins (enoxaparin, dalteparin, etc.), heparin derivatives (fondaparinux, etc.), oral anticoagulants (warfarin, dabigatran etexilate, rivaroxaban, apixaban etc.) except under
specific circumstances of switching oral anticoagulant therapy (see section 4.2) or when UFH is given at doses necessary to maintain an open central venous or arterial catheter (see section 4.5).

- Pregnancy and breast-feeding (see section 4.6).

### 4.4 Special warnings and precautions for use

Lixiana 15 mg is not indicated as monotherapy, as it may result in decreased efficacy. It is only indicated in the process of switching from Lixiana 30 mg (patients with one or more clinical factors for increased exposure; see table 1) to VKA, together with an appropriate VKA dose (see table 2, section 4.2).

**Haemorrhagic risk**

Edoxaban increases the risk of bleeding and can cause serious, potentially fatal bleeding. Lixiana, like other anticoagulants, is recommended to be used with caution in patients with increased risk of bleeding. Lixiana administration should be discontinued if severe haemorrhage occurs (see sections 4.8 and 4.9).

In the clinical studies mucosal bleedings (e.g. epistaxis, gastrointestinal, genitourinary) and anaemia were seen more frequently during long term edoxaban treatment compared with VKA treatment. Thus, in addition to adequate clinical surveillance, laboratory testing of haemoglobin/haematocrit could be of value to detect occult bleeding, as judged to be appropriate.

Several sub-groups of patients, as detailed below, are at increased risk of bleeding. These patients are to be carefully monitored for signs and symptoms of bleeding complications and anaemia after initiation of treatment (see section 4.8). Any unexplained fall in haemoglobin or blood pressure should lead to a search for a bleeding site.

The anticoagulant effect of edoxaban cannot be reliably monitored with standard laboratory testing. A specific anticoagulant reversal agent for edoxaban is not available (see section 4.9).

Haemodialysis does not significantly contribute to edoxaban clearance (see section 5.2).

**Elderly**

The co-administration of Lixiana with ASA in elderly patients should be used cautiously because of a potentially higher bleeding risk (see section 4.5).

**Renal impairment**

The plasma AUC for subjects with mild (CrCL > 50 - 80 mL/min), moderate (CrCL 30 - 50 mL/min) and severe (CrCL < 30 mL/min but not undergoing dialysis) renal impairment was increased by 32%, 74%, and 72%, respectively, relative to subjects with normal renal function (see section 4.2 for dose reduction).

In patients with end stage renal disease or on dialysis, Lixiana is not recommended (see sections 4.2 and 5.2).

**Renal function in NVAF**

A trend towards decreasing efficacy with increasing creatinine clearance was observed for edoxaban compared to well-managed warfarin (see section 5.1). Therefore, edoxaban should only be used in patients with NVAF and high creatinine clearance after a careful evaluation of the individual thromboembolic and bleeding risk.

Assessment of renal function: CrCL should be monitored at the beginning of the treatment in all patients and afterwards when clinically indicated (see section 4.2).

**Hepatic impairment**

Lixiana is not recommended in patients with severe hepatic impairment (see sections 4.2 and 5.2).
Lixiana should be used with caution in patients with mild or moderate hepatic impairment (see section 4.2).

Patients with elevated liver enzymes (ALT/AST > 2 x ULN) or total bilirubin ≥ 1.5 x ULN were excluded in clinical trials. Therefore Lixiana should be used with caution in this population (see sections 4.2 and 5.2). Prior to initiating Lixiana, liver function testing should be performed. Periodic hepatic monitoring is recommended for patients on Lixiana treatment beyond 1 year.

**Discontinuation for surgery and other interventions**
If anticoagulation must be discontinued to reduce the risk of bleeding with surgical or other procedures, Lixiana should be stopped as soon as possible and preferably at least 24 hours before the procedure.

In deciding whether a procedure should be delayed until 24 hours after the last dose of Lixiana, the increased risk of bleeding should be weighed against the urgency of the intervention. Lixiana should be restarted after the surgical or other procedures as soon as adequate haemostasis has been established, noting that the time to onset of the edoxaban anticoagulant therapeutic effect is 1 – 2 hours. If oral medicinal products cannot be taken during or after surgical intervention, consider administering a parenteral anticoagulant and then switch to oral once daily Lixiana (see section 4.2).

**Anticoagulants, antiplatelets, and thrombolytics**
Concomitant use of medicines affecting haemostasis may increase the risk of bleeding. These include acetylsalicylic acid (ASA), P2Y₁₂ platelet inhibitors, other antithrombotic agents, fibrinolytic therapy, and chronic nonsteroidal anti-inflammatory drugs (NSAIDs) (see section 4.5).

**Prosthetic heart valves and moderate to severe mitral stenosis**
Edoxaban has not been studied in patients with mechanical heart valves, in patients during the first 3 months after implantation of a bioprosthetic heart valve, with or without atrial fibrillation, or in patients with moderate to severe mitral stenosis. Therefore, use of edoxaban is not recommended in these patients.

**Haemodynamically unstable PE patients or patients who require thrombolysis or pulmonary embolectomy**
Lixiana is not recommended as an alternative to unfractionated heparin in patients with pulmonary embolism who are haemodynamically unstable or may receive thrombolysis or pulmonary embolectomy since the safety and efficacy of edoxaban have not been established in these clinical situations.

**Patients with active cancer**
Efficacy and safety of edoxaban in the treatment and/or prevention of VTE in patients with active cancer have not been established.

**Laboratory coagulation parameters**
Although treatment with edoxaban does not require routine monitoring, the effect on anticoagulation can be estimated by a calibrated quantitative anti-Factor Xa assay which may help to inform clinical decisions in particular situations as, e.g. overdose and emergency surgery (see also section 5.2).

Edoxaban prolongs standard clotting tests such as prothrombin time (PT), INR, and activated partial thromboplastin time (aPTT) as a result of FXa inhibition. Changes observed in these clotting tests at the expected therapeutic dose are, however, small, subject to a high degree of variability, and not useful in monitoring the anticoagulation effect of edoxaban.
4.5 Interaction with other medicinal products and other forms of interaction

Edoxaban is predominantly absorbed in the upper gastrointestinal (GI) tract. Thus, medicines or disease conditions that increase gastric emptying and gut motility have the possibility of reducing edoxaban dissolution and absorption.

**P-gp inhibitors**

Edoxaban is a substrate for the efflux transporter P-gp. In pharmacokinetic (PK) studies, concomitant administration of edoxaban with the P-gp inhibitors: ciclosporin, dronedarone, erythromycin, ketoconazole, quinidine, or verapamil resulted in increased plasma concentrations of edoxaban. Concomitant use of edoxaban with ciclosporin, dronedarone, erythromycin, or ketoconazole requires dose reduction to 30 mg once daily. Concomitant use of edoxaban with quinidine, verapamil, or amiodarone does not require dose reduction based on clinical data (see section 4.2). The use of edoxaban with other P-gp inhibitors including HIV protease inhibitors has not been studied.

Lixiana 30 mg once daily must be administered during concomitant use with the following P-gp inhibitors:

- **Ciclosporin:** Concurrent administration of a single dose of ciclosporin 500 mg with a single dose of edoxaban 60 mg increased edoxaban AUC and C\text{max} by 73% and 74%, respectively.
- **Dronedarone:** Dronedarone 400 mg twice daily for 7 days with a single concomitant dose of edoxaban 60 mg on Day 5 increased edoxaban AUC and C\text{max} by 85% and 46%, respectively.
- **Erythromycin:** Erythromycin 500 mg four times daily for 8 days with a single concomitant dose of edoxaban 60 mg on Day 7 increased the edoxaban AUC and C\text{max} by 85% and 68%, respectively.
- **Ketoconazole:** Ketoconazole 400 mg once daily for 7 days with a single concomitant dose of edoxaban 60 mg on Day 4, increased edoxaban AUC and C\text{max} by 87% and 89%, respectively.

Lixiana 60 mg once daily is recommended during concomitant use with the following P-gp inhibitors:

- **Quinidine:** Quinidine 300 mg once daily on Days 1 and 4 and three times daily on Days 2 and 3, with a single concomitant dose of edoxaban 60 mg on Day 3, increased edoxaban AUC over 24 hours by 77% and C\text{max} by 85%, respectively.
- **Verapamil:** Verapamil 240 mg once daily for 11 days with a single concomitant dose of edoxaban 60 mg on Day 10 increased the edoxaban AUC and C\text{max} by approximately 53%.
- **Amiodarone:** Co-administration of amiodarone 400 mg once daily with edoxaban 60 mg once daily increased AUC by 40% and C\text{max} by 66%. This was not considered clinically significant. In ENGAGE AF-TIMI 48 study in NVAF, efficacy and safety results were similar for subjects with and without concomitant amiodarone use.

**P-gp inducers**

Co-administration of edoxaban with the P-gp inducer rifampicin led to a decrease in mean edoxaban AUC and a shortened half-life, with possible decreases in its pharmacodynamic effects. The concomitant use of edoxaban with other P-gp inducers (e.g. phenytoin, carbamazepine, phenobarbital or St. John's Wort) may lead to reduced edoxaban plasma concentrations. Edoxaban should be used with caution when co-administered with P-gp inducers.

**P-gp substrates**

- **Digoxin:** Edoxaban 60 mg once daily on days 1 to 14 with coadministration of multiple daily doses of digoxin 0.25 mg twice daily (days 8 and 9) and 0.25 mg once daily (days 10 to 14) increased the C\text{max} of edoxaban by 17%, with no significant effect on AUC or renal clearance at steady state. When the effects of edoxaban on digoxin PK were also examined, the C\text{max} of digoxin increased by approximately 28% and AUC by 7%. This was not considered clinically relevant. No dose modification is necessary when Lixiana is administered with digoxin.
Anticoagulants, antiplatelets, and NSAIDs

Anticoagulants: Co-administration of edoxaban with other anticoagulants is contraindicated due to increased risk of bleeding (see section 4.3).

Acetylsalicylic acid (ASA): Co-administration of ASA (100 mg or 325 mg) and edoxaban increased bleeding time relative to either medicine alone. Co-administration of high dose ASA (325 mg) increased the steady state C_max and AUC of edoxaban by 35% and 32%, respectively. The concomitant chronic use of high dose ASA (325 mg) with edoxaban is not recommended. Concomitant administration of higher doses than 100 mg ASA should only be performed under medical supervision.

In clinical studies concomitant use of ASA (low dose ≤ 100 mg/day), other antiplatelet agents, and thienopyridines was permitted and resulted in approximately a 2-fold increase in major bleeding in comparison with no concomitant use, although to a similar extent in the edoxaban and warfarin groups (see section 4.4). Co-administration of low dose ASA (≤ 100 mg) did not affect the peak or total exposure of edoxaban either after single dose or at steady-state. Edoxaban can be co-administered with low dose ASA (≤ 100 mg/day).

Platelet inhibitors: In ENGAGE AF-TIMI 48 concomitant use of thienopyridines (e.g. clopidogrel) monotherapy was permitted and resulted in increased clinically relevant bleeding although with a lower risk of bleeding on edoxaban compared to warfarin (see section 4.4).

There is very limited experience on the use of edoxaban with dual antiplatelet therapy or fibrinolytic agents.

NSAIDs: Co-administration of naproxen and edoxaban increased bleeding time relative to either medicine alone. Naproxen had no effect on the C_max and AUC of edoxaban. In clinical studies, co-administration of NSAIDs resulted in increased clinically relevant bleeding. Chronic use of NSAIDs with edoxaban is not recommended.

Effect of edoxaban on other medicines
Edoxaban increased the C_max of concomitantly administered digoxin by 28%; however, the AUC was not affected. Edoxaban had no effect on the C_max and AUC of quinidine.

Edoxaban decreased the C_max and AUC of concomitantly administered verapamil by 14% and 16%, respectively.

4.6 Fertility, pregnancy and lactation

Woman of childbearing potential
Women of childbearing potential should avoid becoming pregnant during treatment with edoxaban.

Pregnancy
Safety and efficacy of edoxaban have not been established in pregnant women. Studies in animals have shown reproductive toxicity (see section 5.3). Due to the potential reproductive toxicity, the intrinsic risk of bleeding and the evidence that edoxaban passes the placenta, Lixiana is contraindicated during pregnancy (see section 4.3).

Breast-feeding
Safety and efficacy of edoxaban have not been established in breast-feeding women. Data from animals indicate that edoxaban is secreted into breast milk. Therefore Lixiana is contraindicated during breast-feeding (see section 4.3). A decision must be made whether to discontinue breast-feeding or to discontinue/abstain from therapy.

Fertility
No specific studies with edoxaban in humans have been conducted to evaluate effects on fertility. In a study on male and female fertility in rats no effects were seen (see section 5.3).
4.7 Effects on ability to drive and use machines

Lixiana has no or negligible influence on the ability to drive and use machines.

4.8 Undesirable effects

Summary of the safety profile

The safety of edoxaban has been evaluated in two Phase 3 studies including 21,105 patients with NVAF (ENGAGE AF-TIMI 48 study), and 8,292 patients with VTE (DVT and PE) (Hokusai-VTE study).

The average exposure to edoxaban 60 mg (including 30 mg dose reduced) was 2.5 years among 7,012 patients in ENGAGE AF-TIMI 48 and 251 days among 4,118 patients in Hokusai-VTE.

Adverse reactions were experienced by 2,256 (32.2%) of the patients treated with edoxaban 60 mg (30 mg dose reduced) in the ENGAGE AF-TIMI 48 study and 1,249 (30.3%) in the Hokusai-VTE study.

In both studies, the most common adverse reactions related to bleeding with edoxaban 60 mg based on adjudicated terms included cutaneous soft tissue haemorrhage (up to 5.9%) and epistaxis (up to 4.7%), while vaginal haemorrhage (9.0%) was the most common bleeding-related adverse reaction in Hokusai-VTE only.

Bleeding can occur at any site and may be severe and even fatal (see section 4.4).

Common other adverse reactions for edoxaban were anaemia, rash and abnormal liver function tests.

Tabulated list of adverse reactions

Table 3 provides the list of adverse reactions from the two pivotal Phase 3 studies in patients with VTE (DVT and PE) (Hokusai-VTE study) and AF (ENGAGE AF-TIMI 48 study) combined for both indications. The adverse reactions are classified by System Organ Class and frequency, using the following convention:

Very common (≥ 1/10), Common (≥ 1/100 to < 1/10), Uncommon (≥ 1/1,000 to < 1/100), Rare (≥ 1/10,000 to < 1/1,000), Very rare (< 1/10,000), not known (cannot be estimated from the available data).

Table 3: List of adverse reactions for NVAF and VTE

<table>
<thead>
<tr>
<th>System Organ Class</th>
<th>Frequency</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Blood and lymphatic system disorders</strong></td>
<td></td>
</tr>
<tr>
<td>Anaemia</td>
<td>Common</td>
</tr>
<tr>
<td>Thrombocytopenia</td>
<td>Uncommon</td>
</tr>
<tr>
<td><strong>Immune system disorders</strong></td>
<td></td>
</tr>
<tr>
<td>Hypersensitivity</td>
<td>Uncommon</td>
</tr>
<tr>
<td>Anaphylactic reaction</td>
<td>Rare</td>
</tr>
<tr>
<td>Allergic oedema</td>
<td>Rare</td>
</tr>
<tr>
<td><strong>Nervous system disorders</strong></td>
<td></td>
</tr>
<tr>
<td>Dizziness</td>
<td>Common</td>
</tr>
<tr>
<td>Headache</td>
<td>Common</td>
</tr>
<tr>
<td>Intracranial haemorrhage (ICH)</td>
<td>Uncommon</td>
</tr>
<tr>
<td>Subarachnoid haemorrhage</td>
<td>Rare</td>
</tr>
<tr>
<td><strong>Eye disorders</strong></td>
<td></td>
</tr>
<tr>
<td>Conjunctival/Scleral haemorrhage</td>
<td>Uncommon</td>
</tr>
<tr>
<td>Intraocular haemorrhage</td>
<td>Uncommon</td>
</tr>
<tr>
<td><strong>Cardiac disorders</strong></td>
<td></td>
</tr>
<tr>
<td>Pericardial haemorrhage</td>
<td>Rare</td>
</tr>
<tr>
<td>System Organ Class</td>
<td>Frequency</td>
</tr>
<tr>
<td>--------------------------------------------------------</td>
<td>------------</td>
</tr>
<tr>
<td><strong>Vascular disorders</strong></td>
<td></td>
</tr>
<tr>
<td>Other haemorrhage</td>
<td>Uncommon</td>
</tr>
<tr>
<td><strong>Respiratory, thoracic and mediastinal disorders</strong></td>
<td></td>
</tr>
<tr>
<td>Epistaxis</td>
<td>Common</td>
</tr>
<tr>
<td>Haemoptysis</td>
<td>Uncommon</td>
</tr>
<tr>
<td><strong>Gastrointestinal disorders</strong></td>
<td></td>
</tr>
<tr>
<td>Abdominal pain</td>
<td>Common</td>
</tr>
<tr>
<td>Lower GI haemorrhage</td>
<td>Common</td>
</tr>
<tr>
<td>Upper GI haemorrhage</td>
<td>Common</td>
</tr>
<tr>
<td>Oral/Pharyngeal haemorrhage</td>
<td>Common</td>
</tr>
<tr>
<td>Nausea</td>
<td>Common</td>
</tr>
<tr>
<td>Retroperitoneal haemorrhage</td>
<td>Rare</td>
</tr>
<tr>
<td><strong>Hepatobiliary disorders</strong></td>
<td></td>
</tr>
<tr>
<td>Blood bilirubin increased</td>
<td>Common</td>
</tr>
<tr>
<td>Gammaglutamyltransferase increased</td>
<td>Common</td>
</tr>
<tr>
<td>Blood alkaline phosphatase increased</td>
<td>Uncommon</td>
</tr>
<tr>
<td>Transaminases increased</td>
<td>Uncommon</td>
</tr>
<tr>
<td>Aspartate aminotransferase increased</td>
<td>Uncommon</td>
</tr>
<tr>
<td><strong>Skin and subcutaneous tissue disorders</strong></td>
<td></td>
</tr>
<tr>
<td>Cutaneous soft tissue haemorrhage</td>
<td>Common</td>
</tr>
<tr>
<td>Rash</td>
<td>Common</td>
</tr>
<tr>
<td>Pruritus</td>
<td>Common</td>
</tr>
<tr>
<td>Urticaria</td>
<td>Uncommon</td>
</tr>
<tr>
<td><strong>Musculoskeletal and connective tissue disorders</strong></td>
<td></td>
</tr>
<tr>
<td>Intramuscular haemorrhage (no compartment syndrome)</td>
<td>Rare</td>
</tr>
<tr>
<td>Intra-articular haemorrhage</td>
<td>Rare</td>
</tr>
<tr>
<td><strong>Renal and urinary disorders</strong></td>
<td></td>
</tr>
<tr>
<td>Macroscopic haematuria/urethral haemorrhage</td>
<td>Common</td>
</tr>
<tr>
<td><strong>Reproductive system and breast disorders</strong></td>
<td></td>
</tr>
<tr>
<td>Vaginal haemorrhage</td>
<td>Common</td>
</tr>
<tr>
<td><strong>General disorders and administration site conditions</strong></td>
<td></td>
</tr>
<tr>
<td>Puncture site haemorrhage</td>
<td>Common</td>
</tr>
<tr>
<td><strong>Investigations</strong></td>
<td></td>
</tr>
<tr>
<td>Liver function test abnormal</td>
<td>Common</td>
</tr>
<tr>
<td><strong>Injury, poisoning and procedural complications</strong></td>
<td></td>
</tr>
<tr>
<td>Surgical site haemorrhage</td>
<td>Uncommon</td>
</tr>
<tr>
<td>Subdural haemorrhage</td>
<td>Rare</td>
</tr>
<tr>
<td>Procedural haemorrhage</td>
<td>Rare</td>
</tr>
</tbody>
</table>

1 Reporting rates are based on the female population in clinical trials. Vaginal bleeds were reported commonly in women under the age of 50 years, while it was uncommon in women over the age of 50 years.
Description of selected adverse reactions
Due to the pharmacological mode of action, the use of Lixiana may be associated with an increased risk of occult or overt bleeding from any tissue or organ which may result in post haemorrhagic anaemia. The signs, symptoms, and severity (including fatal outcome) will vary according to the location and degree or extent of the bleeding and/or anaemia (see section 4.9 Management of bleeding). In the clinical studies mucosal bleedings (e.g. epistaxis, gastrointestinal, genitourinary) and anaemia were seen more frequently during long term edoxaban treatment compared with VKA treatment. Thus, in addition to adequate clinical surveillance, laboratory testing of haemoglobin/haematocrit could be of value to detect occult bleeding, as judged to be appropriate. The risk of bleedings may be increased in certain patient groups e.g. those patients with uncontrolled severe arterial hypertension and/or on concomitant treatment affecting haemostasis (see Haemorrhagic risk in section 4.4). Menstrual bleeding may be intensified and/or prolonged. Haemorrhagic complications may present as weakness, paleness, dizziness, headache or unexplained swelling, dyspnoea, and unexplained shock.

Known complications secondary to severe bleeding such as compartment syndrome and renal failure due to hypoperfusion have been reported for Lixiana. Therefore, the possibility of haemorrhage is to be considered in evaluating the condition in any anticoagulated patient.

Reporting of suspected adverse reactions
Reporting suspected adverse reactions after authorisation of the medicinal product is important. It allows continued monitoring of the benefit/risk balance of the medicinal product. Healthcare professionals are asked to report any suspected adverse reactions via the national reporting system listed in Appendix V.

4.9 Overdose

Overdose with edoxaban may lead to haemorrhage. Experience with overdose cases is very limited.

A specific antidote antagonising the pharmacodynamic effect of edoxaban is not available.

Early administration of activated charcoal may be considered in case of edoxaban overdose to reduce absorption. This recommendation is based on standard treatment of drug overdose and data available with similar compounds, as the use of activated charcoal to reduce absorption of edoxaban has not been specifically studied in the edoxaban clinical programme.

Management of bleeding
Should a bleeding complication arise in a patient receiving edoxaban, the next edoxaban administration should be delayed or treatment should be discontinued as appropriate. Edoxaban has a half-life of approximately 10 to 14 hours (see section 5.2). Management should be individualised according to the severity and location of the haemorrhage. Appropriate symptomatic treatment could be used as needed, such as mechanical compression (e.g. for severe epistaxis), surgical haemostasis with bleeding control procedures, fluid replacement and haemodynamic support, blood products (packed red cells or fresh frozen plasma, depending on associated anaemia or coagulopathy) or platelets.

For life-threatening bleeding that cannot be controlled with the measures such as transfusion or haemostasis, the administration of a 4-factor prothrombin complex concentrate (PCC) at 50 IU/kg has been shown to reverse the effects of Lixiana 30 minutes after completing the infusion.

Recombinant factor VIIa (r-FVIIa) can also be considered. However, there is limited clinical experience with the use of this product in individuals receiving edoxaban.

Depending on local availability, a consultation with a coagulation expert should be considered in case of major bleedings.

Protamine sulfate and vitamin K are not expected to affect the anticoagulant activity of edoxaban.
There is no experience with antifibrinolytic agents (tranexamic acid, aminocaproic acid) in individuals receiving edoxaban. There is neither scientific rationale for benefit nor experience with the use of systemic haemostatics (desmopressin, aprotinin) in individuals receiving edoxaban. Due to the high plasma protein binding edoxaban is not expected to be dialysable.

5. PHARMACOLOGICAL PROPERTIES

5.1 Pharmacodynamic properties

Pharmacotherapeutic group: Other antithrombotic agents, ATC code: B01AF03

Mechanism of action
Edoxaban is a highly selective, direct and reversible inhibitor of factor Xa, the serine protease located in the final common pathway of the coagulation cascade. Edoxaban inhibits free factor Xa, and prothrombinase activity. Inhibition of factor Xa in the coagulation cascade reduces thrombin generation, prolongs clotting time and reduces the risk of thrombus formation.

Pharmacodynamic effects
Edoxaban produces rapid onset of pharmacodynamic effects within 1 - 2 hours, which corresponds with peak edoxaban exposure (Cmax). The pharmacodynamic effects measured by anti-factor Xa assay are predictable and correlate with the dose and the concentration of edoxaban. As a result of FXa inhibition, edoxaban also prolongs clotting time in tests such as prothrombin time (PT), and activated partial thromboplastin time (aPTT). Changes observed in these clotting tests are expected at the therapeutic dose, however, these changes are small, subject to a high degree of variability, and not useful in monitoring the anticoagulation effect of edoxaban.

Effects of coagulation markers when switching from rivaroxaban, dabigatran, or apixaban to edoxaban
In clinical pharmacology studies, healthy subjects received rivaroxaban 20 mg once daily, dabigatran 150 mg twice daily, or apixaban 5 mg twice daily, followed by a single dose of edoxaban 60 mg on Day 4. The effect on prothrombin time (PT) and other coagulation biomarkers (e.g. anti-FXa, aPTT) was measured. Following the switch to edoxaban on Day 4 the PT was equivalent to Day 3 of rivaroxaban and apixaban. For dabigatran higher aPTT activity was observed after edoxaban administration with prior dabigatran treatment compared to that after treatment with edoxaban alone. This is considered to be due to the carry-over effect of dabigatran treatment, however, this did not lead to a prolongation of bleeding time.

Based on these data, when switching from these anticoagulants to edoxaban, the first dose of edoxaban can be initiated at the time of the next scheduled dose of the previous anticoagulant (see section 4.2).

Clinical efficacy and safety

Prevention of stroke and systemic embolism
The edoxaban clinical programme for atrial fibrillation was designed to demonstrate the efficacy and safety of two dose groups of edoxaban compared to warfarin for the prevention of stroke and systemic embolism in subjects with nonvalvular atrial fibrillation and at moderate to high risk of stroke and systemic embolic events (SEE).

In the pivotal ENGAGE AF-TIMI 48 study (an event-driven, Phase 3, multi-centre, randomised, double-blind double-dummy parallel-group study), 21,105 subjects, with a mean CHADS2 score of 2.8, were randomised to either edoxaban 30 mg once daily treatment group, or edoxaban 60 mg once daily treatment group or warfarin. Subjects in both edoxaban treatment groups had their dose halved if one or more of the following clinical factors were present: moderate renal impairment (CrCL 30 – 50 mL/min), low body weight (≤ 60 kg) or concomitant use of specific P-gp inhibitors (verapamil, quinidine, dronedarone).
The primary efficacy endpoint was the composite of stroke and SEE. Secondary efficacy endpoints included: Composite of stroke, SEE, and cardiovascular (CV) mortality; major adverse cardiovascular event (MACE), which is the composite of non-fatal MI, non-fatal stroke, non-fatal SEE, and death due to CV cause or bleeding; composite of stroke, SEE, and all-cause mortality.

The median study drug exposure for both the edoxaban 60 mg and 30 mg treatment groups was 2.5 years. The median study follow-up for both the edoxaban 60 mg and 30 mg treatment groups was 2.8 years. The median subject-year exposure was 15,471, and 15,840 for the 60 mg and 30 mg treatment groups, respectively; and the median subject-year follow-up was 19,191 and 19,216 for the 60 mg and 30 mg treatment groups, respectively.

In the warfarin group, the median TTR (time in therapeutic range, INR 2.0 to 3.0) was 68.4%.

The main analysis of efficacy was aimed to show the non-inferiority of edoxaban versus warfarin on first stroke or SEE that occurred during treatment or within 3 days from the last dose taken in the modified intention-to-treat (mITT) population. Edoxaban 60 mg was non-inferior to warfarin for the primary efficacy endpoint of stroke or SEE (upper limit of the 97.5% CI of the HR was below the pre-specified non-inferiority margin of 1.38) (Table 4).

### Table 4: Strokes and Systemic Embolic Events in the ENGAGE AF–TIMI 48 Study - mITT, on-treatment

<table>
<thead>
<tr>
<th>Primary Endpoint</th>
<th>Edoxaban 60 mg (30 mg Dose Reduced) (N = 7,012)</th>
<th>Warfarin (N = 7,012)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>First Stroke/SEE</strong>&lt;sup&gt;a&lt;/sup&gt;</td>
<td></td>
<td></td>
</tr>
<tr>
<td>n</td>
<td>182</td>
<td>232</td>
</tr>
<tr>
<td>Event Rate (%/yr)&lt;sup&gt;b&lt;/sup&gt;</td>
<td>1.18</td>
<td>1.50</td>
</tr>
<tr>
<td>HR (97.5% CI)</td>
<td>0.79 (0.63, 0.99)</td>
<td></td>
</tr>
<tr>
<td>p-value for non-inferiority&lt;sup&gt;c&lt;/sup&gt;</td>
<td>&lt;0.0001</td>
<td></td>
</tr>
<tr>
<td><strong>First Ischaemic Stroke</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>n</td>
<td>135</td>
<td>144</td>
</tr>
<tr>
<td>Event Rate (%/yr)&lt;sup&gt;b&lt;/sup&gt;</td>
<td>0.87</td>
<td>0.93</td>
</tr>
<tr>
<td>HR (95% CI)</td>
<td>0.94 (0.75, 1.19)</td>
<td></td>
</tr>
<tr>
<td><strong>First Haemorrhagic Stroke</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>n</td>
<td>40</td>
<td>76</td>
</tr>
<tr>
<td>Event Rate (%/yr)&lt;sup&gt;b&lt;/sup&gt;</td>
<td>0.26</td>
<td>0.49</td>
</tr>
<tr>
<td>HR (95% CI)</td>
<td>0.53 (0.36, 0.78)</td>
<td></td>
</tr>
<tr>
<td><strong>First SEE</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>n (%/yr)&lt;sup&gt;a&lt;/sup&gt;</td>
<td>8 (0.05)</td>
<td>13 (0.08)</td>
</tr>
<tr>
<td>HR (95% CI)</td>
<td>0.62 (0.26, 1.50)</td>
<td></td>
</tr>
</tbody>
</table>

Abbreviations: HR = Hazard Ratio versus warfarin, CI = Confidence Interval, n = number of events, mITT = modified Intent To Treat, N = number of subjects in mITT population, SEE = Systemic Embolic Event, yr = year.

<sup>a</sup> A subject can be represented in multiple rows.

<sup>b</sup> The event rate (%/yr) is calculated as number of events/subject-year exposure.

<sup>c</sup> The two-sided p-value is based on the non-inferiority margin of 1.38.

During the overall study period in the ITT population (analysis set to show superiority), adjudicated stroke or SEE occurred in 296 subjects in the edoxaban 60 mg group (1.57% per year), and
337 subjects in the warfarin group (1.80% per year). Compared to warfarin-treated subjects, the HR in the edoxaban 60 mg group was 0.87 (99% CI: 0.71, 1.07, p = 0.08 for superiority).

In subgroup analyses, for subjects in the 60 mg treatment group who were dose reduced to 30 mg in the ENGAGE AF-TIMI 48 study (for body weight ≤ 60 kg, moderate renal impairment, or concomitant use of P-gp inhibitors), the event rate was: 2.29% per year for the primary endpoint, compared to the event rate of 2.66% per year for the matching subjects in the warfarin group [HR (95% CI): 0.86 (0.66, 1.13)].

The efficacy results for pre-specified major subgroups (with dose reduction as required), including age, body weight, gender, status of renal function, prior stroke or TIA, diabetes and P-gp inhibitors were generally consistent with the primary efficacy results for the overall population studied in the trial.

The Hazard Ratio (Edoxaban 60 mg vs. warfarin) for the primary endpoint in the centres with a lower average time of INR in the target range (INR TTR) for warfarin was 0.73 – 0.80 for the lowest 3 quartiles (INR TTR ≤ 57.7% to ≤ 73.9%). It was 1.07 in centres with the best control of warfarin therapy (4th quartile with > 73.9% of INR values in the therapeutic range).

There was a statistically significant interaction between the effect of edoxaban versus warfarin on the main study outcome (stroke/SEE) and renal function (p-value 0.0042; mITT, overall study period).

Table 5 shows ischaemic strokes/SEE by creatinine clearance category in NVAF patients in ENGAGE AF-TIMI 48. There is a decreasing event rate at increasing CrCL in both treatment groups.

**Table 5: Number of Ischaemic Strokes/SEE by creatinine clearance category in ENGAGE AF-TIMI 48, mITT Analysis Set Overall Study**

<table>
<thead>
<tr>
<th>CrCL subgroup (mL/min)</th>
<th>Edoxaban 60 mg (N = 7,012)</th>
<th>Warfarin (N = 7,012)</th>
<th>HR (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>n</td>
<td>Number of Events</td>
<td>Event rate (%/year)</td>
</tr>
<tr>
<td>≥ 30 to ≤ 50</td>
<td>1,302</td>
<td>63</td>
<td>1.89</td>
</tr>
<tr>
<td>&gt; 50 to ≤ 70</td>
<td>2,093</td>
<td>85</td>
<td>1.51</td>
</tr>
<tr>
<td>&gt; 70 to ≤ 90</td>
<td>1,661</td>
<td>45</td>
<td>0.99</td>
</tr>
<tr>
<td>&gt; 90 to ≤ 110</td>
<td>927</td>
<td>27</td>
<td>1.08</td>
</tr>
<tr>
<td>&gt; 110 to ≤ 130</td>
<td>497</td>
<td>14</td>
<td>1.01</td>
</tr>
<tr>
<td>&gt; 130</td>
<td>462</td>
<td>10</td>
<td>0.78</td>
</tr>
</tbody>
</table>

Abbreviations: N = number of subjects; mITT population overall study period; n = number of patients in subgroup
*HR not computed if number of events < 5 in one treatment group.

Within renal function subgroups, results for the secondary efficacy endpoints were consistent with those for the primary endpoint.

Superiority testing was performed on the ITT Overall Study Period. Stroke and SEE occurred in fewer subjects in the edoxaban 60 mg treatment group than in the warfarin group (1.57% and 1.80% per year, respectively), with a HR of 0.87 (99% CI: 0.71, 1.07, p = 0.0807 for superiority).
The pre-specified composite endpoints for the comparison of the edoxaban 60 mg treatment group to warfarin for stroke, SEE, and CV mortality HR (99% CI) was 0.87 (0.76, 0.99), MACE 0.89 (0.78, 1.00), and stroke, SEE, and all-cause mortality 0.90 (0.80, 1.01).

The results for all-cause mortality (adjudicated deaths) in the ENGAGE AF-TIMI 48 study were 769 (3.99% per year) for subjects taking edoxaban 60 mg (30 mg dose reduced) as opposed to 836 (4.35% per year) for warfarin [HR (95% CI): 0.91 (0.83, 1.01)].

All-cause mortality (adjudicated deaths) per renal subgroups (edoxaban vs. warfarin): CrCL 30 to ≤ 50 mL/min [HR (95% CI): 0.81 (0.68, 0.97)]; CrCL > 50 to < 80 mL/min [HR (95% CI): 0.87 (0.75, 1.02)]; CrCL ≥ 80 mL/min [HR (95% CI): 1.15 (0.95, 1.40)].

Edoxaban 60 mg (30 mg dose reduced) resulted in a lower rate of cardiovascular mortality compared to warfarin [HR (95% CI): 0.86 (0.77, 0.97)].

Adjudicated efficacy cardiovascular mortality per renal subgroups (edoxaban vs. warfarin): CrCL 30 to ≤ 50 mL/min [HR (95% CI): 0.80 (0.65, 0.99)]; CrCL > 50 to < 80 mL/min [HR (95% CI): 0.75 (0.62, 0.90)]; CrCL ≥ 80 mL/min [HR (95% CI): 1.16 (0.92, 1.46)].

Safety in patients with NVAF in ENGAGE AF-TIMI 48

The primary safety endpoint was major bleeding.

There was a significant risk reduction in favour of the edoxaban 60 mg treatment group compared with the warfarin group in major bleeding (2.75%, and 3.43% per year, respectively) [HR (95% CI): 0.80 (0.71, 0.91); p = 0.0009], ICH (0.39%, and 0.85% per year, respectively) [HR (95% CI): 0.47 (0.34, 0.63); p < 0.0001], and other types of bleeding (Table 6).

The reduction in fatal bleeds was also significant for the edoxaban 60 mg treatment group compared with the warfarin group [HR (95% CI): 0.55 (0.36, 0.84); p = 0.0059 for superiority], primarily because of the reduction in fatal ICH bleeds [HR (95% CI): 0.58 (0.35, 0.95); p = 0.0312].
Table 6: Bleeding Events in ENGAGE AF-TIMI 48 Study - Safety Analysis On-Treatment

<table>
<thead>
<tr>
<th>Major Bleeding</th>
<th>Edoxaban 60 mg (30 mg Dose Reduced) (N = 7,012)</th>
<th>Warfarin (N = 7,012)</th>
</tr>
</thead>
<tbody>
<tr>
<td>n</td>
<td>418</td>
<td>524</td>
</tr>
<tr>
<td>Event rate (%/yr)&lt;sup&gt;a&lt;/sup&gt;</td>
<td>2.75</td>
<td>3.43</td>
</tr>
<tr>
<td>HR (95% CI)</td>
<td>0.80 (0.71, 0.91)</td>
<td>0.0009</td>
</tr>
<tr>
<td>n</td>
<td>61</td>
<td>132</td>
</tr>
<tr>
<td>Event rate (%/yr)&lt;sup&gt;a&lt;/sup&gt;</td>
<td>0.39</td>
<td>0.85</td>
</tr>
<tr>
<td>HR (95% CI)</td>
<td>0.47 (0.34, 0.63)</td>
<td></td>
</tr>
<tr>
<td>Fatal Bleeding</td>
<td></td>
<td></td>
</tr>
<tr>
<td>n</td>
<td>32</td>
<td>59</td>
</tr>
<tr>
<td>Event rate (%/yr)&lt;sup&gt;a&lt;/sup&gt;</td>
<td>0.21</td>
<td>0.38</td>
</tr>
<tr>
<td>CRNM Bleeding</td>
<td></td>
<td></td>
</tr>
<tr>
<td>n</td>
<td>1,214</td>
<td>1,396</td>
</tr>
<tr>
<td>Event rate (%/yr)&lt;sup&gt;a&lt;/sup&gt;</td>
<td>8.67</td>
<td>10.15</td>
</tr>
<tr>
<td>HR (95% CI)</td>
<td>0.86 (0.80, 0.93)</td>
<td></td>
</tr>
<tr>
<td>Any Confirmed Bleeding&lt;sup&gt;c&lt;/sup&gt;</td>
<td></td>
<td></td>
</tr>
<tr>
<td>n</td>
<td>1,865</td>
<td>2,114</td>
</tr>
<tr>
<td>Event rate (%/yr)&lt;sup&gt;a&lt;/sup&gt;</td>
<td>14.15</td>
<td>16.40</td>
</tr>
<tr>
<td>HR (95% CI)</td>
<td>0.87 (0.82, 0.92)</td>
<td></td>
</tr>
</tbody>
</table>

Abbreviations: ICH = Intracranial Haemorrhage, HR = Hazard Ratio versus warfarin, CI = Confidence Interval, CRNM = Clinically Relevant Non-Major, n = number of subjects with events, N = number of subjects in Safety population, yr = year.

<sup>a</sup> The event rate (%/yr) is calculated as number of events/subject-year exposure.

<sup>b</sup> ICH includes primary haemorrhagic stroke, subarachnoid haemorrhage, epidural/subdural haemorrhage, and ischaemic stroke with major haemorrhagic conversion. All ICHs reported on the Adjudicated Cerebrovascular and Non-Intracranial bleed eCRF forms confirmed by the adjudicators are included in ICH counts.

<sup>c</sup> 'Any Confirmed Bleeding' includes those that the adjudicator defined as clinically overt.

Note: A subject can be included in multiple sub-categories if he/she had an event for those categories. The first event of each category is included in the analysis.

Tables 7, 8 and 9 show major, fatal and intracranial bleedings, respectively, by creatinine clearance category in NVAF patients in ENGAGE AF-TIMI 48. There is a decreasing event rate at increasing CrCL in both treatment groups.
### Table 7: Number of Major Bleeding Events by creatinine clearance category in ENGAGE AF-TIMI 48, Safety Analysis On-Treatment\(^a\)

<table>
<thead>
<tr>
<th>CrCL subgroup (mL/min)</th>
<th>Edoxaban 60 mg (N = 7,012)</th>
<th>Warfarin (N = 7,012)</th>
<th>HR (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>n</td>
<td>Number of Events</td>
<td>Event rate (%/year)</td>
</tr>
<tr>
<td>≥ 30 to ≤ 50</td>
<td>1,302</td>
<td>96</td>
<td>3.91</td>
</tr>
<tr>
<td>&gt; 50 to ≤ 70</td>
<td>2,093</td>
<td>148</td>
<td>3.31</td>
</tr>
<tr>
<td>&gt; 70 to ≤ 90</td>
<td>1,661</td>
<td>108</td>
<td>2.88</td>
</tr>
<tr>
<td>&gt; 90 to ≤ 110</td>
<td>927</td>
<td>29</td>
<td>1.33</td>
</tr>
<tr>
<td>&gt; 110 to ≤ 130</td>
<td>497</td>
<td>20</td>
<td>1.70</td>
</tr>
<tr>
<td>&gt; 130</td>
<td>462</td>
<td>13</td>
<td>1.18</td>
</tr>
</tbody>
</table>

### Table 8: Number of Fatal Bleeding Events by creatinine clearance category in ENGAGE AF-TIMI 48, Safety Analysis On-Treatment\(^a\)

<table>
<thead>
<tr>
<th>CrCL subgroup (mL/min)</th>
<th>Edoxaban 60 mg (N = 7,012)</th>
<th>Warfarin (N = 7,012)</th>
<th>HR (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>n</td>
<td>Number of Events</td>
<td>Event rate (%/year)</td>
</tr>
<tr>
<td>≥ 30 to ≤ 50</td>
<td>1,302</td>
<td>9</td>
<td>0.36</td>
</tr>
<tr>
<td>&gt; 50 to ≤ 70</td>
<td>2,093</td>
<td>8</td>
<td>0.18</td>
</tr>
<tr>
<td>&gt; 70 to ≤ 90</td>
<td>1,661</td>
<td>10</td>
<td>0.26</td>
</tr>
<tr>
<td>&gt; 90 to ≤ 110</td>
<td>927</td>
<td>2</td>
<td>0.09</td>
</tr>
<tr>
<td>&gt; 110 to ≤ 130</td>
<td>497</td>
<td>1</td>
<td>0.08</td>
</tr>
<tr>
<td>&gt; 130</td>
<td>462</td>
<td>2</td>
<td>0.18</td>
</tr>
</tbody>
</table>

### Table 9: Number of Intracranial Bleeding Events by creatinine clearance category in ENGAGE AF-TIMI 48, Safety Analysis On-Treatment\(^a\)

<table>
<thead>
<tr>
<th>CrCL subgroup (mL/min)</th>
<th>Edoxaban 60 mg (N = 7,012)</th>
<th>Warfarin (N = 7,012)</th>
<th>HR (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>n</td>
<td>Number of Events</td>
<td>Event rate (%/year)</td>
</tr>
<tr>
<td>≥ 30 to ≤ 50</td>
<td>1,302</td>
<td>16</td>
<td>0.64</td>
</tr>
<tr>
<td>&gt; 50 to ≤ 70</td>
<td>2,093</td>
<td>19</td>
<td>0.42</td>
</tr>
<tr>
<td>&gt; 70 to ≤ 90</td>
<td>1,661</td>
<td>17</td>
<td>0.44</td>
</tr>
<tr>
<td>&gt; 90 to ≤ 110</td>
<td>927</td>
<td>5</td>
<td>0.23</td>
</tr>
<tr>
<td>&gt; 110 to ≤ 130</td>
<td>497</td>
<td>2</td>
<td>0.17</td>
</tr>
<tr>
<td>&gt; 130</td>
<td>462</td>
<td>1</td>
<td>0.09</td>
</tr>
</tbody>
</table>

Abbreviations: N = number of subjects; mITT population overall study period; n = number of patients in subgroup

*HR not computed if number of events < 5 in one treatment group.

\(^a\) On-Treatment: Time from first dose of study drug to last dose plus 3 days.
In subgroup analyses, for subjects in the 60 mg treatment group who were dose reduced to 30 mg in the ENGAGE AF-TIMI 48 study for body weight ≤ 60 kg, moderate renal impairment, or concomitant use of P-gp inhibitors, 104 (3.05% per year) of edoxaban 30 mg dose reduced subjects and 166 (4.85% per year) of warfarin dose reduced subjects had a major bleeding event [HR (95% CI): 0.63 (0.50, 0.81)].

In the ENGAGE AF-TIMI 48 study there was a significant improvement in Net Clinical Outcome (First Stroke, SEE, Major Bleed, or All-Cause Mortality; mITT population, overall study period) in favour of edoxaban, HR (95% CI): 0.89 (0.83, 0.96); p = 0.0024, when edoxaban 60 mg treatment group was compared to warfarin.

Treatment of DVT, treatment of PE and the prevention of recurrent DVT and PE (VTE)
The edoxaban clinical programme for VTE was designed to demonstrate the efficacy and safety of edoxaban in the treatment of DVT and PE, and the prevention of recurrent DVT and PE.

In the pivotal Hokusai-VTE study, 8,292 subjects were randomised to receive initial heparin therapy (enoxaparin or unfractionated heparin) followed by edoxaban 60 mg once daily or the comparator. In the comparator arm, subjects received initial heparin therapy concurrently with warfarin, titrated to a target INR of 2.0 to 3.0, followed by warfarin alone. The treatment duration was from 3 months up to 12 months, determined by the investigator based on the patient’s clinical features.

The majority of edoxaban treated patients were Caucasians (69.6%) and Asians (21.0%), 3.8% were Black, 5.3% were categorised as Other race.

The duration of therapy was at least 3 months for 3,718 (91.6%) edoxaban subjects versus 3,727 (91.4%) of warfarin subjects; at least 6 months for 3,495 (86.1%) of edoxaban subjects versus 3,491 (85.6%) of warfarin subjects; and 12 months for 1,643 (40.5%) edoxaban subjects versus 1,659 (40.4%) of warfarin subjects.

The primary efficacy endpoint was the recurrence of symptomatic VTE, defined as the composite of recurrent symptomatic DVT, non-fatal symptomatic PE and fatal PE in subjects during the 12-month study period. Secondary efficacy outcomes included the composite clinical outcome of recurrent VTE and all-cause mortality.

Edoxaban 30 mg once daily was used for subjects with one or more of the following clinical factors: moderate renal impairment (CrCL 30 - 50 mL/min); body weight ≤ 60 kg; concomitant use of specific P-gp inhibitors.

In the Hokusai-VTE study (Table 10) edoxaban was demonstrated to be non-inferior to warfarin for the primary efficacy outcome, recurrent VTE, which occurred in 130 of 4,118 subjects (3.2%) in the edoxaban group versus 146 of 4,122 subjects (3.5%) in the warfarin group [HR (95% CI): 0.89 (0.70, 1.13); p < 0.0001 for non-inferiority]. In the warfarin group, the median TTR (time in therapeutic range, INR 2.0 to 3.0) was 65.6%. For subjects presenting with PE (with or without DVT), 47 (2.8%) of edoxaban and 65 (3.9%) of warfarin subjects had a recurrent VTE [HR (95% CI): 0.73 (0.50, 1.06)].
Table 10: Efficacy Results from the Hokusai-VTE Study - mITT population, overall study period

<table>
<thead>
<tr>
<th>Primary endpoint(^a)</th>
<th>Edoxaban 60 mg (30 mg Dose Reduced) (N = 4,118)</th>
<th>Warfarin (N = 4,122)</th>
<th>Edoxaban vs Warfarin HR (95% CI)(^b) p-value(^c)</th>
</tr>
</thead>
<tbody>
<tr>
<td>All subjects with symptomatic recurrent VTE(^c), n (%)</td>
<td>130 (3.2)</td>
<td>146 (3.5)</td>
<td>0.89 (0.70, 1.13) p-value &lt; 0.0001 (non-inferiority)</td>
</tr>
<tr>
<td>PE with or without DVT</td>
<td>73 (1.8)</td>
<td>83 (2.0)</td>
<td></td>
</tr>
<tr>
<td>Fatal PE or Death where PE cannot be ruled out</td>
<td>24 (0.6)</td>
<td>24 (0.6)</td>
<td></td>
</tr>
<tr>
<td>Non-fatal PE</td>
<td>49 (1.2)</td>
<td>59 (1.4)</td>
<td></td>
</tr>
<tr>
<td>DVT only</td>
<td>57 (1.4)</td>
<td>63 (1.5)</td>
<td></td>
</tr>
</tbody>
</table>

Abbreviations: CI = Confidence Interval; DVT = deep vein thrombosis; mITT = modified intent-to-treat; HR = Hazard Ratio vs. warfarin; n = number of subjects with events; N = number of subjects in mITT population; PE = pulmonary embolism; VTE = venous thromboembolic events.

\(^a\) The primary efficacy endpoint is adjudicated symptomatic recurrent VTE (i.e., the composite endpoint of DVT, non-fatal PE, and fatal PE).

\(^b\) The HR, two-sided CI are based on the Cox proportional hazards regression model including treatment and the following randomisation stratification factors as covariates: presenting diagnosis (PE with or without DVT, DVT only), baseline risk factors (temporary factors, all others), and the need for 30 mg edoxaban/edoxaban placebo dose at randomisation (yes/no).

\(^c\) The p-value is for the pre-defined non-inferiority margin of 1.5.

For the subjects who were dose reduced to 30 mg (predominantly low body weight or renal function) 15 (2.1%) edoxaban and 22 (3.1%) of warfarin subjects had a recurrent VTE [HR (95% CI): 0.69 (0.36, 1.34)].

The secondary composite endpoint of recurrent VTE and all-cause mortality occurred in 138 subjects (3.4%) in the edoxaban group and 158 subjects (3.9%) in the warfarin group [HR (95% CI): 0.87 (0.70, 1.10)].

The results for all-cause mortality (adjudicated deaths) in Hokusai-VTE were 136 (3.3%) for subjects taking edoxaban 60 mg (30 mg dose reduced) as opposed to 130 (3.2%) for warfarin.

In a pre-specified subgroup analysis of PE subjects 447 (30.6%) and 483 (32.2%) of edoxaban and warfarin treated subjects, respectively, were identified as having PE and NT-proBNP ≥ 500 pg/mL. The primary efficacy outcome occurred in 14 (3.1%) and 30 (6.2%) of edoxaban and warfarin subjects, respectively [HR (95% CI): 0.50 (0.26, 0.94)].

The efficacy results for pre-specified major subgroups (with dose reduction as required), including age, body weight, gender and status of renal function were consistent with the primary efficacy results for the overall population studied in the trial.

**Safety in patients with VTE (DVT and PE) in Hokusai-VTE**

The primary safety endpoint was clinically relevant bleeding (major or clinically relevant non-major).

Table 11 summarises adjudicated bleeding events for the safety analysis set on-treatment period.
There was a significant risk reduction in favour of edoxaban compared with warfarin for the primary safety endpoint of clinically relevant bleeding, a composite of major bleeding or clinically relevant non-major bleeding (CRNM), which occurred in 349 of 4,118 subjects (8.5%) in the edoxaban group and in 423 of 4,122 subjects (10.3%) in the warfarin group \[HR (95\% CI): 0.81 (0.71, 0.94); p = 0.004 for superiority\].

Table 11: Bleeding Events in Hokusai-VTE Study - Safety Analysis On-Treatment Period\(^a\)

<table>
<thead>
<tr>
<th>Clinically Relevant Bleeding (Major and CRNM)(^b), n (%)</th>
<th>Edoxaban 60 mg (30 mg Dose Reduced) (N = 4,118)</th>
<th>Warfarin (N = 4,122)</th>
</tr>
</thead>
<tbody>
<tr>
<td>n</td>
<td>349 (8.5)</td>
<td>423 (10.3)</td>
</tr>
<tr>
<td>HR (95% CI)</td>
<td>0.81 (0.71, 0.94)</td>
<td></td>
</tr>
<tr>
<td>p-value</td>
<td>0.004 (for superiority)</td>
<td></td>
</tr>
<tr>
<td>Major Bleeding n (%)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>n</td>
<td>56 (1.4)</td>
<td>66 (1.6)</td>
</tr>
<tr>
<td>HR (95% CI)</td>
<td>0.84 (0.59, 1.21)</td>
<td></td>
</tr>
<tr>
<td>ICH fatal</td>
<td>0</td>
<td>6 (0.1)</td>
</tr>
<tr>
<td>ICH non-fatal</td>
<td>5 (0.1)</td>
<td>12 (0.3)</td>
</tr>
<tr>
<td>CRNM Bleeding</td>
<td></td>
<td></td>
</tr>
<tr>
<td>n</td>
<td>298 (7.2)</td>
<td>368 (8.9)</td>
</tr>
<tr>
<td>HR (95% CI)</td>
<td>0.80 (0.68, 0.93)</td>
<td></td>
</tr>
<tr>
<td>All Bleeding</td>
<td></td>
<td></td>
</tr>
<tr>
<td>n</td>
<td>895 (21.7)</td>
<td>1,056 (25.6)</td>
</tr>
<tr>
<td>HR (95% CI)</td>
<td>0.82 (0.75, 0.90)</td>
<td></td>
</tr>
</tbody>
</table>

Abbreviations: ICH = Intracranial Haemorrhage, HR = Hazard Ratio vs. warfarin; CI = Confidence Interval; N = number of subjects in safety population; n = number of events; CRNM = clinically relevant non-major

\(^a\) On-Treatment Period: Time from first dose of study drug to last dose plus 3 days.

\(^b\) Primary Safety Endpoint: Clinically relevant bleeding (composite of major and clinically relevant non-major bleeding).

In subgroup analyses, for subjects who were dose reduced to 30 mg in the Hokusai-VTE study for body weight ≤ 60 kg, moderate renal impairment, or concomitant use of P-gp inhibitors, 58 (7.9%) of edoxaban 30 mg dose reduced subjects and 92 (12.8%) of warfarin subjects had a major bleeding or CRNM event \[HR (95\%): 0.62 (0.44, 0.86)\].

In the Hokusai-VTE study the Net Clinical Outcome (Recurrent VTE, Major Bleed, or All-Cause Mortality; mITT population, overall study period) HR (95\% CI) was 1.00 (0.85, 1.18) when edoxaban was compared to warfarin.

Patients undergoing cardioversion

A multicentre, prospective, randomised, open-label study with blinded endpoint evaluation (ENSURE-AF) was conducted which randomised 2199 subjects (oral anticoagulant naïve and pre-treated) with non-valvular atrial fibrillation scheduled for cardioversion, to compare edoxaban 60 mg once daily with enoxaparin/warfarin to maintain a therapeutic INR of 2.0 to 3.0 (randomised 1:1), mean TTR on warfarin was 70.8%. A total of 2149 subjects were treated with either edoxaban (N = 1067) or enoxaparin/warfarin (N = 1082). Subjects in the edoxaban treatment group received 30 mg once daily
if one or more of the following clinical factors were present: moderate renal impairment (CrCL 30 – 50 mL/min), low body weight (≤ 60 kg) or concomitant use of specific P-gp inhibitors. The majority of subjects in the edoxaban and warfarin groups had cardioversion performed (83.7% and 78.9%, respectively) or were auto-converted (6.6% and 8.6%, respectively). TEE-guided (within 3 days of initiation) or conventional cardioversion (at least 21 days of pre-treatment) was employed. Subjects were maintained on treatment for 28 days post cardioversion.

The primary efficacy outcome consisted of a composite of all stroke, SEE, MI and CV mortality. A total of 5 (0.5%, 95% CI 0.15% - 1.06%) events occurred in subjects in the edoxaban group (N = 1095) and 11 (1.0%, 95% CI 0.50% - 1.78%) events in the warfarin group (N = 1104); OR 0.46 (95% CI 0.12 - 1.43); ITT analysis set overall study period with mean duration of 66 days.

The primary safety outcome was a composite of major and CRNM bleeding. A total of 16 (1.5%, 95% CI 0.86% - 2.42%) events occurred in subjects in the edoxaban (N = 1067) group and 11 (1.0%, 95% CI 0.51% - 1.81%) events in the warfarin (N = 1082) group; OR 1.48 (95% CI 0.64 - 3.55); safety analysis set on-treatment period.

This exploratory study showed low rates of major and CRNM bleeding and thromboembolism in the two treatment groups in the setting of cardioversion.

Paediatric population
The European Medicines Agency has deferred the obligation to submit the results of studies with edoxaban in one or more subsets of the paediatric population in prevention of arterial thrombosis, treatment of thromboembolism and prevention of thromboembolism (see section 4.2 for information on paediatric use).

5.2 Pharmacokinetic properties
Absorption
Edoxaban is absorbed with peak plasma concentrations within 1 - 2 hours. The absolute bioavailability is approximately 62%. Food increases peak exposure to a varying extent, but has minimal effect on total exposure. Edoxaban was administered with or without food in the ENGAGE AF-TIMI 48 and the Hokusai-VTE studies. Edoxaban is poorly soluble at pH of 6.0 or higher. Co-administration of proton-pump inhibitors had no relevant impact on edoxaban exposure.

Distribution
Disposition is biphasic. The volume of distribution is 107 (19.9) L mean (SD). In vitro plasma protein binding is approximately 55%. There is no clinically relevant accumulation of edoxaban (accumulation ratio 1.14) with once daily dosing. Steady state concentrations are achieved within 3 days.

Biotransformation
Unchanged edoxaban is the predominant form in plasma. Edoxaban is metabolised via hydrolysis (mediated by carboxylesterase 1), conjugation or oxidation by CYP3A4/5 (< 10%). Edoxaban has three active metabolites, the predominant metabolite (M-4), formed by hydrolysis, is active and reaches less than 10% of the exposure of the parent compound in healthy subjects. Exposure to the other metabolites is less than 5%. Edoxaban is a substrate for the efflux transporter P-glycoprotein (P-gp), but not a substrate for uptake transporters such as organic anion transporter polypeptide OATP1B1, organic anion transporters OAT1 or OAT3 or organic cation transporter OCT2. Its active metabolite is a substrate for OATP1B1.

Elimination
In healthy subjects, the total clearance is estimated as 22 (± 3) L/hour; 50% is renally cleared (11 L/hour). Renal clearance accounts for approximately 35% of the administered dose. Metabolism and biliary/intestinal excretion account for the remaining clearance. The t½ for oral administration is 10 - 14 hours.
**Linearity/non-linearity**
Edoxaban displays approximately dose-proportional pharmacokinetics for doses of 15 mg to 60 mg in healthy subjects.

**Special populations**

*Elderly*
After taking renal function and body weight into account, age had no additional clinically significant effect on edoxaban pharmacokinetics in a population pharmacokinetic analysis of the pivotal Phase 3 study in NVAF (ENGAGE AF-TIMI 48).

*Gender*
After accounting for body weight, gender had no additional clinically significant effect on edoxaban pharmacokinetics in a population pharmacokinetic analysis of the Phase 3 study in NVAF (ENGAGE AF-TIMI 48).

*Ethnic origin*
In a population pharmacokinetic analysis of the ENGAGE AF-TIMI 48 study, peak and total exposure in Asian patients and non-Asian patients were comparable.

**Renal impairment**
The plasma AUCs for subjects with mild (CrCL > 50 - 80 mL/min), moderate (CrCL 30 - 50 mL/min) and severe (CrCL < 30 mL/min but not undergoing dialysis) renal impairment were increased by 32%, 74%, and 72%, respectively, relative to subjects with normal renal function. In patients with renal impairment the metabolite profile changes and a higher quantity of active metabolites are formed. There is a linear correlation between edoxaban plasma concentration and anti-FXa activity regardless of renal function.
Subjects with ESRD undergoing peritoneal dialysis had 93% higher total exposure compared with healthy subjects.
Population PK modeling indicates that exposure approximately doubles in patients with severe renal impairment (CrCL 15 – 29 mL/min) relative to patients with normal renal function.

*Anti-FXa activity by CrCL category*
Table 12 below shows the edoxaban anti-Factor Xa activity by CrCL category for each indication.
Table 12: Edoxaban Anti-FXa activity by creatinine clearance

<table>
<thead>
<tr>
<th>Edoxaban Dose</th>
<th>CrCL (mL/min)</th>
<th>Edoxaban Anti-FXa activity post-dose (IU/mL) (^1)</th>
<th>Edoxaban Anti-FXa activity pre-dose (IU/mL) (^2)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Prevention of stroke and systemic embolism: NVAF</td>
<td></td>
<td></td>
</tr>
<tr>
<td>30 mg QD</td>
<td>≥ 30 to ≤ 50</td>
<td>2.92 [0.33 – 5.88]</td>
<td>0.53 [0.11 – 2.06]</td>
</tr>
<tr>
<td></td>
<td>&gt; 50 to ≤ 70</td>
<td>4.52 [0.38 – 7.64]</td>
<td>0.83 [0.16 – 2.61]</td>
</tr>
<tr>
<td></td>
<td>&gt; 70 to ≤ 90</td>
<td>4.12 [0.19 – 7.55]</td>
<td>0.68 [0.05 – 2.33]</td>
</tr>
<tr>
<td></td>
<td>&gt; 90 to ≤ 110</td>
<td>3.82 [0.36 – 7.39]</td>
<td>0.60 [0.14 – 3.57]</td>
</tr>
<tr>
<td></td>
<td>&gt; 110 to ≤ 130</td>
<td>3.16 [0.28 – 6.71]</td>
<td>0.41 [0.15 – 1.51]</td>
</tr>
<tr>
<td></td>
<td>&gt; 130</td>
<td>2.76 [0.12 – 6.10]</td>
<td>0.45 [0.00 – 3.10]</td>
</tr>
<tr>
<td></td>
<td>Treatment of DVT, treatment of PE and prevention of recurrent DVT and PE (VTE)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>30 mg QD</td>
<td>≥ 30 to ≤ 50</td>
<td>2.21 [0.14 – 4.47]</td>
<td>0.22 [0.00 – 1.09]</td>
</tr>
<tr>
<td></td>
<td>&gt; 50 to ≤ 70</td>
<td>3.42 [0.19 – 6.13]</td>
<td>0.34 [0.00 – 3.10]</td>
</tr>
<tr>
<td></td>
<td>&gt; 70 to ≤ 90</td>
<td>2.97 [0.24 – 5.82]</td>
<td>0.24 [0.00 – 1.77]</td>
</tr>
<tr>
<td></td>
<td>&gt; 90 to ≤ 110</td>
<td>2.82 [0.14 – 5.31]</td>
<td>0.20 [0.00 – 2.52]</td>
</tr>
<tr>
<td></td>
<td>&gt; 110 to ≤ 130</td>
<td>2.64 [0.13 – 5.57]</td>
<td>0.17 [0.00 – 1.86]</td>
</tr>
<tr>
<td></td>
<td>&gt; 130</td>
<td>2.39 [0.10 – 4.92]</td>
<td>0.13 [0.00 – 2.43]</td>
</tr>
</tbody>
</table>

\(^*\)Dose reduction to 30 mg for low body weight ≤ 60 kg or specific concomitant P-glycoprotein (P-gp) inhibitors

\(^1\) Post-dose is equivalent to C\(_{\text{max}}\) (post-dose samples were drawn 1 – 3 hours after edoxaban administration)

\(^2\) Pre-dose is equivalent to C\(_{\text{min}}\)

Although treatment with edoxaban does not require routine monitoring, the effect on anticoagulation can be estimated by a calibrated quantitative anti-Factor Xa assay which may be useful in exceptional situations where knowledge of edoxaban exposure may help to inform clinical decisions, e.g. overdose and emergency surgery (see also section 4.4).

**Haemodialysis**
A 4 hour haemodialysis session reduced total edoxaban exposures by less than 9%.

**Hepatic impairment**
Patients with mild or moderate hepatic impairment exhibited comparable pharmacokinetics and pharmacodynamics to their matched healthy control group. Edoxaban has not been studied in patients with severe hepatic impairment (see section 4.2).
**Body weight**

In a population pharmacokinetic analysis of the ENGAGE AF-TIMI 48 study in NVAF, $C_{\text{max}}$ and $AUC$ in patients with median low body weight (55 kg) were increased by 40% and 13%, respectively, as compared with patients with median high body weight (84 kg). In Phase 3 clinical studies (both NVAF and VTE indications) patients with body weight ≤ 60 kg had a 50% edoxaban dose reduction and had similar efficacy and less bleeding when compared to warfarin.

**Pharmacokinetic/pharmacodynamic relationship(s)**

PT, INR, aPTT and Anti-factor Xa correlate linearly with edoxaban concentrations.

### 5.3 Preclinical safety data

Non-clinical data reveal no special hazard for humans based on conventional studies of safety pharmacology, repeated dose toxicity, genotoxicity, carcinogenic potential, or phototoxicity.

**Reproductive toxicology**

Edoxaban showed vaginal haemorrhage at higher doses in rats and rabbits but had no effects in the reproductive performance of parent rats.

In rats, no effects on male or female fertility were seen.

In animal reproduction studies, rabbits showed increased incidence of gallbladder variations at a dosage of 200 mg/kg which is approximately 65 times the maximum recommended human dose (MRHD) of 60 mg/day based on total body surface area in mg/m². Increased post-implantation pregnancy losses occurred in rats at 300 mg/kg/day (approximately 49 times the MRHD) and in rabbits at 200 mg/kg/day (approximately 65 times the MRHD) respectively.

Edoxaban was excreted in the breast milk of lactating rats.

**Environmental Risk Assessment (ERA)**

The active substance edoxaban tosilate is persistent in the environment (for instructions on disposal see section 6.6).

### 6. PHARMACEUTICAL PARTICULARS

#### 6.1 List of excipients

**Tablet core:**
- Mannitol (E421)
- Pregelatinised starch
- Crospovidone
- Hydroxypropylcellulose
- Magnesium stearate (E470b)

**Film-coat:**
- Hypromellose (E464)
- Macrogol 8000
- Titanium dioxide (E171)
- Talc
- Carnauba wax
- Iron oxide yellow (E172)
- Iron oxide red (E172)

#### 6.2 Incompatibilities

Not applicable.
6.3 Shelf life

5 years

6.4 Special precautions for storage

This medicinal product does not require any special storage conditions.

6.5 Nature and contents of container

PVC/Aluminium blisters. Cartons of 10 film-coated tablets.
PVC/Aluminium perforated unit dose blisters of 10 x 1 film-coated tablets.

Not all pack sizes may be marketed.

6.6 Special precautions for disposal

Any unused medicinal product or waste material should be disposed of in accordance with local requirements.

7. MARKETING AUTHORISATION HOLDER

Daiichi Sankyo Europe GmbH
Zielstattstrasse 48
81379 Munich
Germany

8. MARKETING AUTHORIZATION NUMBER(S)

EU/1/15/993/001
EU/1/15/993/016

9. DATE OF FIRST AUTHORIZATION/RENEWAL OF THE AUTHORIZATION

Date of first authorisation: 19 June 2015

10. DATE OF REVISION OF THE TEXT

Detailed information on this medicinal product is available on the website of the European Medicines Agency http://www.ema.europa.eu.
This medicinal product is subject to additional monitoring. This will allow quick identification of new safety information. Healthcare professionals are asked to report any suspected adverse reactions. See section 4.8 for how to report adverse reactions.

1. **NAME OF THE MEDICINAL PRODUCT**

Lixiana 30 mg film-coated tablets

2. **QUALITATIVE AND QUANTITATIVE COMPOSITION**

Each film-coated tablet contains 30 mg edoxaban (as tosilate).

For the full list of excipients, see section 6.1.

3. **PHARMACEUTICAL FORM**

Film-coated tablet.
Pink, round-shaped film-coated tablets (8.5 mm diameter) debossed with “DSC L30”.

4. **CLINICAL PARTICULARS**

4.1 **Therapeutic indications**

Prevention of stroke and systemic embolism in adult patients with nonvalvular atrial fibrillation (NVAF) with one or more risk factors, such as congestive heart failure, hypertension, age ≥ 75 years, diabetes mellitus, prior stroke or transient ischaemic attack (TIA).

Treatment of deep vein thrombosis (DVT) and pulmonary embolism (PE), and prevention of recurrent DVT and PE in adults (see section 4.4 for haemodynamically unstable PE patients).

4.2 **Posology and method of administration**

**Posology**

*Prevention of stroke and systemic embolism*
The recommended dose is 60 mg edoxaban once daily.
Therapy with edoxaban in NVAF patients should be continued long term.

*Treatment of DVT, treatment of PE and prevention of recurrent DVT and PE (VTE)*
The recommended dose is 60 mg edoxaban once daily following initial use of parenteral anticoagulant for at least 5 days (see section 5.1). Edoxaban and initial parenteral anticoagulant should not be administered simultaneously.
The duration of therapy for treatment of DVT and PE (venous thromboembolism, VTE), and prevention of recurrent VTE should be individualised after careful assessment of the treatment benefit against the risk for bleeding (see section 4.4). Short duration of therapy (at least 3 months) should be based on transient risk factors (e.g. recent surgery, trauma, immobilisation) and longer durations should be based on permanent risk factors or idiopathic DVT or PE.
For NVAF and VTE the recommended dose is 30 mg edoxaban once daily in patients with one or more of the following clinical factors:

- Moderate or severe renal impairment (creatinine clearance (CrCL) 15 - 50 mL/min)
- Low body weight ≤ 60 kg
- Concomitant use of the following P-glycoprotein (P-gp) inhibitors: ciclosporin, dronedarone, erythromycin, or ketoconazole.

### Table 1: Summary of posology in NVAF and VTE (DVT and PE)

<table>
<thead>
<tr>
<th>Summary Guide for Dosing</th>
<th>60 mg once daily</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Recommended dose</strong></td>
<td></td>
</tr>
<tr>
<td><strong>Dose recommendation for patients with one or more of the following clinical factors:</strong></td>
<td></td>
</tr>
<tr>
<td>Renal Impairment</td>
<td></td>
</tr>
<tr>
<td>Moderate or severe (CrCL 15 – 50 mL/min)</td>
<td>30 mg once daily</td>
</tr>
<tr>
<td>Low Body Weight</td>
<td></td>
</tr>
<tr>
<td>≤ 60 kg</td>
<td></td>
</tr>
<tr>
<td>P-gp Inhibitors</td>
<td></td>
</tr>
<tr>
<td>Ciclosporin, dronedarone, erythromycin, ketoconazole</td>
<td></td>
</tr>
</tbody>
</table>

**Missed dose**
If a dose of Lixiana is missed, the dose should be taken immediately and then be continued the following day with the once-daily intake as recommended. The patient should not take double the prescribed dose on the same day to make up for a missed dose.

**Switching to and from Lixiana**
Continued anticoagulant therapy is important in patients with NVAF and VTE. There may be situations that warrant a change in anticoagulation therapy (Table 2).

### Table 2: Switching

<table>
<thead>
<tr>
<th>Switching to Lixiana</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>From</strong></td>
</tr>
<tr>
<td>Vitamin K Antagonist (VKA)</td>
</tr>
<tr>
<td>Oral anticoagulants other than VKA</td>
</tr>
<tr>
<td>• dabigatran</td>
</tr>
<tr>
<td>• rivaroxaban</td>
</tr>
<tr>
<td>• apixaban</td>
</tr>
<tr>
<td>Parenteral anticoagulants</td>
</tr>
</tbody>
</table>
## Switching from Lixiana

<table>
<thead>
<tr>
<th>From</th>
<th>To</th>
<th>Recommendation</th>
</tr>
</thead>
</table>
| Lixiana                       | Vitamin K Antagonist (VKA)| There is a potential for inadequate anticoagulation during the transition from Lixiana to VKA. Continuous adequate anticoagulation should be ensured during any transition to an alternate anticoagulant.  

**Oral option:** For patients currently on a 60 mg dose, administer a Lixiana dose of 30 mg once daily together with an appropriate VKA dose.  

For patients currently on a 30 mg dose (for one or more of the following clinical factors: moderate to severe renal impairment (CrCL 15 – 50 mL/min), low body weight, or use with certain P-gp inhibitors), administer a Lixiana dose of 15 mg once daily together with an appropriate VKA dose.  

Patients should not take a loading dose of VKA in order to promptly achieve a stable INR between 2 and 3. It is recommended to take into account the maintenance dose of VKA and if the patient was previously taking a VKA or to use valid INR driven VKA treatment algorithm, in accordance with local practice.  

Once an INR ≥ 2.0 is achieved, Lixiana should be discontinued. Most patients (85%) should be able to achieve an INR ≥ 2.0 within 14 days of concomitant administration of Lixiana and VKA. After 14 days it is recommended that Lixiana is discontinued and the VKA continued to be titrated to achieve an INR between 2 and 3.  

It is recommended that during the first 14 days of concomitant therapy the INR is measured at least 3 times just prior to taking the daily dose of Lixiana to minimise the influence of Lixiana on INR measurements. Concomitant Lixiana and VKA can increase the INR post Lixiana dose by up to 46%.  

**Parenteral option:** Discontinue Lixiana and administer a parenteral anticoagulant and VKA at the time of the next scheduled Lixiana dose. Once a stable INR of ≥ 2.0 is achieved, the parenteral anticoagulant should be discontinued and the VKA continued.  

| Lixiana                       | Oral anticoagulants other than VKA | Discontinue Lixiana and start the non-VKA anticoagulant at the time of the next scheduled dose of Lixiana. |
Switching from Lixiana

<table>
<thead>
<tr>
<th>From</th>
<th>To</th>
<th>Recommendation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lixiana</td>
<td>Parenteral</td>
<td>These agents should not be administered simultaneously. Discontinue Lixiana and start the parenteral anticoagulant at the time of the next scheduled dose of Lixiana.</td>
</tr>
</tbody>
</table>

**Special populations**

**Assessment of renal function:**
- Renal function should be assessed in all patients by calculating the creatinine clearance (CrCL) prior to initiation of treatment with Lixiana to exclude patients with end stage renal disease (i.e. CrCL < 15 mL/min), to use the correct Lixiana dose in patients with CrCL 15 – 50 mL/min (30 mg once daily), in patients with CrCL > 50 mL/min (60 mg once daily) and when deciding on the use of Lixiana in patients with increased creatinine clearance (see section 4.4).
- Renal function should also be assessed when a change in renal function is suspected during treatment (e.g. hypovolaemia, dehydration, and in case of concomitant use of certain medicinal products).

The method used to estimate renal function (CrCL in mL/min) during the clinical development of Lixiana was the Cockcroft-Gault method. The formula is as follows:

For creatinine in µmol/L:

\[
1.23 \times \frac{(140-\text{age [years]}) \times \text{weight [kg]} (\times 0.85 \text{ if female})}{\text{serum creatinine [µmol/L]}}
\]

For creatinine in mg/dL:

\[
\frac{(140-\text{age [years]}) \times \text{weight [kg]} (\times 0.85 \text{ if female})}{72 \times \text{serum creatinine [mg/dL]}}
\]

This method is recommended when assessing patients’ CrCL prior to and during Lixiana treatment.

**Renal impairment**

In patients with mild renal impairment (CrCL > 50 – 80 mL/min), the recommended dose is 60 mg Lixiana once daily.

In patients with moderate or severe renal impairment (CrCL 15 – 50 mL/min), the recommended dose is 30 mg Lixiana once daily (see section 5.2).

In patients with end stage renal disease (ESRD) (CrCL < 15 mL/min) or on dialysis, the use of Lixiana is not recommended (see sections 4.4 and 5.2).

**Hepatic impairment**

Lixiana is contraindicated in patients with hepatic disease associated with coagulopathy and clinically relevant bleeding risk (see section 4.3).

In patients with severe hepatic impairment Lixiana is not recommended (see sections 4.4 and 5.2).

In patients with mild to moderate hepatic impairment the recommended dose is 60 mg Lixiana once daily (see section 5.2). Lixiana should be used with caution in patients with mild to moderate hepatic impairment (see section 4.4).
Patients with elevated liver enzymes (ALT/AST > 2 x ULN) or total bilirubin ≥ 1.5 x ULN were excluded in clinical trials. Therefore Lixiana should be used with caution in this population (see sections 4.4 and 5.2). Prior to initiating Lixiana, liver function testing should be performed.

Body weight
For patients with body weight ≤ 60 kg, the recommended dose is 30 mg Lixiana once daily (see section 5.2).

Elderly
No dose reduction is required (see section 5.2).

Gender
No dose reduction is required (see section 5.2).

Concomitant use of Lixiana with P-glycoprotein (P-gp) inhibitors
In patients concomitantly taking Lixiana and the following P-gp inhibitors: ciclosporin, dronedarone, erythromycin, or ketoconazole, the recommended dose is 30 mg Lixiana once daily (see section 4.5). No dose reduction is required for concomitant use of amiodarone, quinidine or verapamil (see section 4.5).
The use of Lixiana with other P-gp inhibitors including HIV protease inhibitors has not been studied.

Paediatric population
The safety and efficacy of Lixiana in children and adolescents less than 18 years of age have not been established. No data are available.

Patients undergoing cardioversion
Lixiana can be initiated or continued in patients who may require cardioversion. For transoesophageal echocardiogram (TEE) guided cardioversion in patients not previously treated with anticoagulants, Lixiana treatment should be started at least 2 hours before cardioversion to ensure adequate anticoagulation (see sections 5.1 and 5.2). Cardioversion should be performed no later than 12 hours after the dose of Lixiana on the day of the procedure.

**For all patients undergoing cardioversion:** Confirmation should be sought prior to cardioversion that the patient has taken Lixiana as prescribed. Decisions on initiation and duration of treatment should follow established guidelines for anticoagulant treatment in patients undergoing cardioversion.

Method of administration
For oral use.
Lixiana can be taken with or without food (see section 5.2).

4.3 Contraindications

- Hypersensitivity to the active substance or to any of the excipients listed in section 6.1.
- Clinically significant active bleeding.
- Hepatic disease associated with coagulopathy and clinically relevant bleeding risk.
- Lesion or condition, if considered to be a significant risk for major bleeding. This may include current or recent gastrointestinal ulceration, presence of malignant neoplasms at high risk of bleeding, recent brain or spinal injury, recent brain, spinal or ophthalmic surgery, recent intracranial haemorrhage, known or suspected oesophageal varices, arteriovenous malformations, vascular aneurysms or major intraspinal or intracerebral vascular abnormalities.
- Uncontrolled severe hypertension.
- Concomitant treatment with any other anticoagulants e.g. unfractionated heparin (UFH), low molecular weight heparins (enoxaparin, dalteparin, etc.), heparin derivatives (fondaparinux, etc.), oral anticoagulants (warfarin, dabigatran etexilate, rivaroxaban, apixaban etc.) except under
specific circumstances of switching oral anticoagulant therapy (see section 4.2) or when UFH is given at doses necessary to maintain an open central venous or arterial catheter (see section 4.5).

- Pregnancy and breast-feeding (see section 4.6).

### 4.4 Special warnings and precautions for use

Lixiana 15 mg is not indicated as monotherapy, as it may result in decreased efficacy. It is only indicated in the process of switching from Lixiana 30 mg (patients with one or more clinical factors for increased exposure; see table 1) to VKA, together with an appropriate VKA dose (see table 2, section 4.2).

**Haemorrhagic risk**

Edoxaban increases the risk of bleeding and can cause serious, potentially fatal bleeding. Lixiana, like other anticoagulants, is recommended to be used with caution in patients with increased risk of bleeding. Lixiana administration should be discontinued if severe haemorrhage occurs (see sections 4.8 and 4.9).

In the clinical studies mucosal bleedings (e.g. epistaxis, gastrointestinal, genitourinary) and anaemia were seen more frequently during long term edoxaban treatment compared with VKA treatment. Thus, in addition to adequate clinical surveillance, laboratory testing of haemoglobin/haematocrit could be of value to detect occult bleeding, as judged to be appropriate.

Several sub-groups of patients, as detailed below, are at increased risk of bleeding. These patients are to be carefully monitored for signs and symptoms of bleeding complications and anaemia after initiation of treatment (see section 4.8). Any unexplained fall in haemoglobin or blood pressure should lead to a search for a bleeding site.

The anticoagulant effect of edoxaban cannot be reliably monitored with standard laboratory testing. A specific anticoagulant reversal agent for edoxaban is not available (see section 4.9).

Haemodialysis does not significantly contribute to edoxaban clearance (see section 5.2).

**Elderly**

The co-administration of Lixiana with ASA in elderly patients should be used cautiously because of a potentially higher bleeding risk (see section 4.5).

**Renal impairment**

The plasma AUC for subjects with mild (CrCL > 50 - 80 mL/min), moderate (CrCL 30 - 50 mL/min) and severe (CrCL < 30 mL/min but not undergoing dialysis) renal impairment was increased by 32%, 74%, and 72%, respectively, relative to subjects with normal renal function (see section 4.2 for dose reduction).

In patients with end stage renal disease or on dialysis, Lixiana is not recommended (see sections 4.2 and 5.2).

**Renal function in NVAF**

A trend towards decreasing efficacy with increasing creatinine clearance was observed for edoxaban compared to well-managed warfarin (see section 5.1). Therefore, edoxaban should only be used in patients with NVAF and high creatinine clearance after a careful evaluation of the individual thromboembolic and bleeding risk.

Assessment of renal function: CrCL should be monitored at the beginning of the treatment in all patients and afterwards when clinically indicated (see section 4.2).

**Hepatic impairment**

Lixiana is not recommended in patients with severe hepatic impairment (see sections 4.2 and 5.2).
Lixiana should be used with caution in patients with mild or moderate hepatic impairment (see section 4.2).

Patients with elevated liver enzymes (ALT/AST > 2 x ULN) or total bilirubin ≥ 1.5 x ULN were excluded in clinical trials. Therefore Lixiana should be used with caution in this population (see sections 4.2 and 5.2). Prior to initiating Lixiana, liver function testing should be performed. Periodic hepatic monitoring is recommended for patients on Lixiana treatment beyond 1 year.

**Discontinuation for surgery and other interventions**

If anticoagulation must be discontinued to reduce the risk of bleeding with surgical or other procedures, Lixiana should be stopped as soon as possible and preferably at least 24 hours before the procedure.

In deciding whether a procedure should be delayed until 24 hours after the last dose of Lixiana, the increased risk of bleeding should be weighed against the urgency of the intervention. Lixiana should be restarted after the surgical or other procedures as soon as adequate haemostasis has been established, noting that the time to onset of the edoxaban anticoagulant therapeutic effect is 1 – 2 hours. If oral medicinal products cannot be taken during or after surgical intervention, consider administering a parenteral anticoagulant and then switch to oral once daily Lixiana (see section 4.2).

**Anticoagulants, antiplatelets, and thrombolytics**

Concomitant use of medicines affecting haemostasis may increase the risk of bleeding. These include acetylsalicylic acid (ASA), P2Y12 platelet inhibitors, other antithrombotic agents, fibrinolytic therapy, and chronic nonsteroidal anti-inflammatory drugs (NSAIDs) (see section 4.5).

**Prosthetic heart valves and moderate to severe mitral stenosis**

Edoxaban has not been studied in patients with mechanical heart valves, in patients during the first 3 months after implantation of a bioprosthetic heart valve, with or without atrial fibrillation, or in patients with moderate to severe mitral stenosis. Therefore, use of edoxaban is not recommended in these patients.

**Haemodynamically unstable PE patients or patients who require thrombolysis or pulmonary embolectomy**

Lixiana is not recommended as an alternative to unfractionated heparin in patients with pulmonary embolism who are haemodynamically unstable or may receive thrombolysis or pulmonary embolectomy since the safety and efficacy of edoxaban have not been established in these clinical situations.

**Patients with active cancer**

Efficacy and safety of edoxaban in the treatment and/or prevention of VTE in patients with active cancer have not been established.

**Laboratory coagulation parameters**

Although treatment with edoxaban does not require routine monitoring, the effect on anticoagulation can be estimated by a calibrated quantitative anti-Factor Xa assay which may help to inform clinical decisions in particular situations as, e.g. overdose and emergency surgery (see also section 5.2).

Edoxaban prolongs standard clotting tests such as prothrombin time (PT), INR, and activated partial thromboplastin time (aPTT) as a result of FXa inhibition. Changes observed in these clotting tests at the expected therapeutic dose are, however, small, subject to a high degree of variability, and not useful in monitoring the anticoagulation effect of edoxaban.
4.5 Interaction with other medicinal products and other forms of interaction

Edoxaban is predominantly absorbed in the upper gastrointestinal (GI) tract. Thus, medicines or disease conditions that increase gastric emptying and gut motility have the possibility of reducing edoxaban dissolution and absorption.

P-gp inhibitors

Edoxaban is a substrate for the efflux transporter P-gp. In pharmacokinetic (PK) studies, concomitant administration of edoxaban with the P-gp inhibitors: ciclosporin, dronedarone, erythromycin, ketoconazole, quinine, or verapamil resulted in increased plasma concentrations of edoxaban. Concomitant use of edoxaban with ciclosporin, dronedarone, erythromycin, or ketoconazole requires dose reduction to 30 mg once daily. Concomitant use of edoxaban with quinine, verapamil, or amiodarone does not require dose reduction based on clinical data (see section 4.2). The use of edoxaban with other P-gp inhibitors including HIV protease inhibitors has not been studied.

Lixiana 30 mg once daily must be administered during concomitant use with the following P-gp inhibitors:

- **Ciclosporin**: Concurrent administration of a single dose of ciclosporin 500 mg with a single dose of edoxaban 60 mg increased edoxaban AUC and C_max by 73% and 74%, respectively.
- **Dronedarone**: Dronedarone 400 mg twice daily for 7 days with a single concomitant dose of edoxaban 60 mg on Day 5 increased edoxaban AUC and C_max by 85% and 46%, respectively.
- **Erythromycin**: Erythromycin 500 mg four times daily for 8 days with a single concomitant dose of edoxaban 60 mg on Day 7 increased the edoxaban AUC and C_max by 85% and 68%, respectively.
- **Ketoconazole**: Ketoconazole 400 mg once daily for 7 days with a single concomitant dose of edoxaban 60 mg on Day 4, increased edoxaban AUC and C_max by 87% and 89%, respectively.

Lixiana 60 mg once daily is recommended during concomitant use with the following P-gp inhibitors:

- **Quinidine**: Quinidine 300 mg once daily on Days 1 and 4 and three times daily on Days 2 and 3, with a single concomitant dose of edoxaban 60 mg on Day 3, increased edoxaban AUC over 24 hours by 77% and C_max by 85%, respectively.
- **Verapamil**: Verapamil 240 mg once daily for 11 days with a single concomitant dose of edoxaban 60 mg on Day 10 increased the edoxaban AUC and C_max by approximately 53%.
- **Amiodarone**: Co-administration of amiodarone 400 mg once daily with edoxaban 60 mg once daily increased AUC by 40% and C_max by 66%. This was not considered clinically significant. In ENGAGE AF-TIMI 48 study in NVAF, efficacy and safety results were similar for subjects with and without concomitant amiodarone use.

P-gp inducers

Co-administration of edoxaban with the P-gp inducer rifampicin led to a decrease in mean edoxaban AUC and a shortened half-life, with possible decreases in its pharmacodynamic effects. The concomitant use of edoxaban with other P-gp inducers (e.g. phenytoin, carbamazepine, phenobarbital or St. John's Wort) may lead to reduced edoxaban plasma concentrations. Edoxaban should be used with caution when co-administered with P-gp inducers.

P-gp substrates

**Digoxin**: Edoxaban 60 mg once daily on days 1 to 14 with co-administration of multiple daily doses of digoxin 0.25 mg twice daily (days 8 and 9) and 0.25 mg once daily (days 10 to 14) increased the C_max of edoxaban by 17%, with no significant effect on AUC or renal clearance at steady state. When the effects of edoxaban on digoxin PK were also examined, the C_max of digoxin increased by approximately 28% and AUC by 7%. This was not considered clinically relevant. No dose modification is necessary when Lixiana is administered with digoxin.
Anticoagulants, antiplatelets, and NSAIDs

Anticoagulants: Co-administration of edoxaban with other anticoagulants is contraindicated due to increased risk of bleeding (see section 4.3).

Acetylsalicylic acid (ASA): Co-administration of ASA (100 mg or 325 mg) and edoxaban increased bleeding time relative to either medicine alone. Co-administration of high dose ASA (325 mg) increased the steady state $C_{\text{max}}$ and AUC of edoxaban by 35% and 32%, respectively. The concomitant chronic use of high dose ASA (325 mg) with edoxaban is not recommended. Concomitant administration of higher doses than 100 mg ASA should only be performed under medical supervision.

In clinical studies concomitant use of ASA (low dose $\leq 100$ mg/day), other antiplatelet agents, and thienopyridines was permitted and resulted in approximately a 2-fold increase in major bleeding in comparison with no concomitant use, although to a similar extent in the edoxaban and warfarin groups (see section 4.4). Co-administration of low dose ASA ($\leq 100$ mg) did not affect the peak or total exposure of edoxaban either after single dose or at steady-state.

Edoxaban can be co-administered with low dose ASA ($\leq 100$ mg/day).

Platelet inhibitors: In ENGAGE AF-TIMI 48 concomitant use of thienopyridines (e.g. clopidogrel) monotherapy was permitted and resulted in increased clinically relevant bleeding although with a lower risk of bleeding on edoxaban compared to warfarin (see section 4.4).

There is very limited experience on the use of edoxaban with dual antiplatelet therapy or fibrinolytic agents.

NSAIDs: Co-administration of naproxen and edoxaban increased bleeding time relative to either medicine alone. Naproxen had no effect on the $C_{\text{max}}$ and AUC of edoxaban. In clinical studies, co-administration of NSAIDs resulted in increased clinically relevant bleeding. Chronic use of NSAIDs with edoxaban is not recommended.

Effect of edoxaban on other medicines

Edoxaban increased the $C_{\text{max}}$ of concomitantly administered digoxin by 28%; however, the AUC was not affected. Edoxaban had no effect on the $C_{\text{max}}$ and AUC of quinidine.

Edoxaban decreased the $C_{\text{max}}$ and AUC of concomitantly administered verapamil by 14% and 16%, respectively.

4.6 Fertility, pregnancy and lactation

Woman of childbearing potential

Women of childbearing potential should avoid becoming pregnant during treatment with edoxaban.

Pregnancy

Safety and efficacy of edoxaban have not been established in pregnant women. Studies in animals have shown reproductive toxicity (see section 5.3). Due to the potential reproductive toxicity, the intrinsic risk of bleeding and the evidence that edoxaban passes the placenta, Lixiana is contraindicated during pregnancy (see section 4.3).

Breast-feeding

Safety and efficacy of edoxaban have not been established in breast-feeding women. Data from animals indicate that edoxaban is secreted into breast milk. Therefore Lixiana is contraindicated during breast-feeding (see section 4.3). A decision must be made whether to discontinue breast-feeding or to discontinue/abstain from therapy.

Fertility

No specific studies with edoxaban in humans have been conducted to evaluate effects on fertility. In a study on male and female fertility in rats no effects were seen (see section 5.3).
4.7 Effects on ability to drive and use machines
Lixiana has no or negligible influence on the ability to drive and use machines.

4.8 Undesirable effects

Summary of the safety profile
The safety of edoxaban has been evaluated in two Phase 3 studies including 21,105 patients with NVAF (ENGAGE AF-TIMI 48 study), and 8,292 patients with VTE (DVT and PE) (Hokusai-VTE study).

The average exposure to edoxaban 60 mg (including 30 mg dose reduced) was 2.5 years among 7,012 patients in ENGAGE AF-TIMI 48 and 251 days among 4,118 patients in Hokusai-VTE. Adverse reactions were experienced by 2,256 (32.2%) of the patients treated with edoxaban 60 mg (30 mg dose reduced) in the ENGAGE AF-TIMI 48 study and 1,249 (30.3%) in the Hokusai-VTE study.

In both studies, the most common adverse reactions related to bleeding with edoxaban 60 mg based on adjudicated terms included cutaneous soft tissue haemorrhage (up to 5.9%) and epistaxis (up to 4.7%), while vaginal haemorrhage (9.0%) was the most common bleeding-related adverse reaction in Hokusai-VTE only.

Bleeding can occur at any site and may be severe and even fatal (see section 4.4). Common other adverse reactions for edoxaban were anaemia, rash and abnormal liver function tests.

Tabulated list of adverse reactions
Table 3 provides the list of adverse reactions from the two pivotal Phase 3 studies in patients with VTE (DVT and PE) (Hokusai-VTE study) and AF (ENGAGE AF-TIMI 48 study) combined for both indications. The adverse reactions are classified by System Organ Class and frequency, using the following convention:
Very common (≥ 1/10), Common (≥ 1/100 to < 1/10), Uncommon (≥ 1/1,000 to < 1/100), Rare (≥ 1/10,000 to < 1/1,000), Very rare (< 1/10,000), not known (cannot be estimated from the available data).

Table 3: List of adverse reactions for NVAF and VTE

<table>
<thead>
<tr>
<th>System Organ Class</th>
<th>Frequency</th>
</tr>
</thead>
<tbody>
<tr>
<td>Blood and lymphatic system disorders</td>
<td></td>
</tr>
<tr>
<td>Anaemia</td>
<td>Common</td>
</tr>
<tr>
<td>Thrombocytopenia</td>
<td>Uncommon</td>
</tr>
<tr>
<td>Immune system disorders</td>
<td></td>
</tr>
<tr>
<td>Hypersensitivity</td>
<td>Uncommon</td>
</tr>
<tr>
<td>Anaphylactic reaction</td>
<td>Rare</td>
</tr>
<tr>
<td>Allergic oedema</td>
<td>Rare</td>
</tr>
<tr>
<td>Nervous system disorders</td>
<td></td>
</tr>
<tr>
<td>Dizziness</td>
<td>Common</td>
</tr>
<tr>
<td>Headache</td>
<td>Common</td>
</tr>
<tr>
<td>Intracranial haemorrhage (ICH)</td>
<td>Uncommon</td>
</tr>
<tr>
<td>Subarachnoid haemorrhage</td>
<td>Rare</td>
</tr>
<tr>
<td>Eye disorders</td>
<td></td>
</tr>
<tr>
<td>Conjunctival/Scleral haemorrhage</td>
<td>Uncommon</td>
</tr>
<tr>
<td>Intraocular haemorrhage</td>
<td>Uncommon</td>
</tr>
<tr>
<td>Cardiac disorders</td>
<td></td>
</tr>
<tr>
<td>Pericardial haemorrhage</td>
<td>Rare</td>
</tr>
<tr>
<td>System Organ Class</td>
<td>Frequency</td>
</tr>
<tr>
<td>--------------------------------------------------------</td>
<td>------------</td>
</tr>
<tr>
<td><strong>Vascular disorders</strong></td>
<td></td>
</tr>
<tr>
<td>Other haemorrhage</td>
<td>Uncommon</td>
</tr>
<tr>
<td><strong>Respiratory, thoracic and mediastinal disorders</strong></td>
<td></td>
</tr>
<tr>
<td>Epistaxis</td>
<td>Common</td>
</tr>
<tr>
<td>Haemoptysis</td>
<td>Uncommon</td>
</tr>
<tr>
<td><strong>Gastrointestinal disorders</strong></td>
<td></td>
</tr>
<tr>
<td>Abdominal pain</td>
<td>Common</td>
</tr>
<tr>
<td>Lower GI haemorrhage</td>
<td>Common</td>
</tr>
<tr>
<td>Upper GI haemorrhage</td>
<td>Common</td>
</tr>
<tr>
<td>Oral/Pharyngeal haemorrhage</td>
<td>Common</td>
</tr>
<tr>
<td>Nausea</td>
<td>Common</td>
</tr>
<tr>
<td>Retroperitoneal haemorrhage</td>
<td>Rare</td>
</tr>
<tr>
<td><strong>Hepatobiliary disorders</strong></td>
<td></td>
</tr>
<tr>
<td>Blood bilirubin increased</td>
<td>Common</td>
</tr>
<tr>
<td>Gammaglutamyltransferase increased</td>
<td>Common</td>
</tr>
<tr>
<td>Blood alkaline phosphatase increased</td>
<td>Uncommon</td>
</tr>
<tr>
<td>Transaminases increased</td>
<td>Uncommon</td>
</tr>
<tr>
<td>Aspartate aminotransferase increased</td>
<td>Uncommon</td>
</tr>
<tr>
<td><strong>Skin and subcutaneous tissue disorders</strong></td>
<td></td>
</tr>
<tr>
<td>Cutaneous soft tissue haemorrhage</td>
<td>Common</td>
</tr>
<tr>
<td>Rash</td>
<td>Common</td>
</tr>
<tr>
<td>Pruritus</td>
<td>Common</td>
</tr>
<tr>
<td>Urticaria</td>
<td>Uncommon</td>
</tr>
<tr>
<td><strong>Musculoskeletal and connective tissue disorders</strong></td>
<td></td>
</tr>
<tr>
<td>Intramuscular haemorrhage (no compartment syndrome)</td>
<td>Rare</td>
</tr>
<tr>
<td>Intra-articular haemorrhage</td>
<td>Rare</td>
</tr>
<tr>
<td><strong>Renal and urinary disorders</strong></td>
<td></td>
</tr>
<tr>
<td>Macroscopic haematuria/urethral haemorrhage</td>
<td>Common</td>
</tr>
<tr>
<td><strong>Reproductive system and breast disorders</strong></td>
<td></td>
</tr>
<tr>
<td>Vaginal haemorrhage</td>
<td>Common</td>
</tr>
<tr>
<td><strong>General disorders and administration site conditions</strong></td>
<td></td>
</tr>
<tr>
<td>Puncture site haemorrhage</td>
<td>Common</td>
</tr>
<tr>
<td><strong>Investigations</strong></td>
<td></td>
</tr>
<tr>
<td>Liver function test abnormal</td>
<td>Common</td>
</tr>
<tr>
<td><strong>Injury, poisoning and procedural complications</strong></td>
<td></td>
</tr>
<tr>
<td>Surgical site haemorrhage</td>
<td>Uncommon</td>
</tr>
<tr>
<td>Subdural haemorrhage</td>
<td>Rare</td>
</tr>
<tr>
<td>Procedural haemorrhage</td>
<td>Rare</td>
</tr>
</tbody>
</table>

1 Reporting rates are based on the female population in clinical trials. Vaginal bleeds were reported commonly in women under the age of 50 years, while it was uncommon in women over the age of 50 years.
Description of selected adverse reactions
Due to the pharmacological mode of action, the use of Lixiana may be associated with an increased risk of occult or overt bleeding from any tissue or organ which may result in post haemorrhagic anaemia. The signs, symptoms, and severity (including fatal outcome) will vary according to the location and degree or extent of the bleeding and/or anaemia (see section 4.9 Management of bleeding). In the clinical studies mucosal bleedings (e.g. epistaxis, gastrointestinal, genitourinary) and anaemia were seen more frequently during long term edoxaban treatment compared with VKA treatment. Thus, in addition to adequate clinical surveillance, laboratory testing of haemoglobin/haematocrit could be of value to detect occult bleeding, as judged to be appropriate. The risk of bleedings may be increased in certain patient groups e.g. those patients with uncontrolled severe arterial hypertension and/or on concomitant treatment affecting haemostasis (see Haemorrhagic risk in section 4.4). Menstrual bleeding may be intensified and/or prolonged. Haemorrhagic complications may present as weakness, paleness, dizziness, headache or unexplained swelling, dyspnoea, and unexplained shock.
Known complications secondary to severe bleeding such as compartment syndrome and renal failure due to hypoperfusion have been reported for Lixiana. Therefore, the possibility of haemorrhage is to be considered in evaluating the condition in any anticoagulated patient.

Reporting of suspected adverse reactions
Reporting suspected adverse reactions after authorisation of the medicinal product is important. It allows continued monitoring of the benefit/risk balance of the medicinal product. Healthcare professionals are asked to report any suspected adverse reactions via the national reporting system listed in Appendix V.

4.9 Overdose
Overdose with edoxaban may lead to haemorrhage. Experience with overdose cases is very limited.
A specific antidote antagonising the pharmacodynamic effect of edoxaban is not available.

Early administration of activated charcoal may be considered in case of edoxaban overdose to reduce absorption. This recommendation is based on standard treatment of drug overdose and data available with similar compounds, as the use of activated charcoal to reduce absorption of edoxaban has not been specifically studied in the edoxaban clinical programme.

Management of bleeding
Should a bleeding complication arise in a patient receiving edoxaban, the next edoxaban administration should be delayed or treatment should be discontinued as appropriate. Edoxaban has a half-life of approximately 10 to 14 hours (see section 5.2). Management should be individualised according to the severity and location of the haemorrhage. Appropriate symptomatic treatment could be used as needed, such as mechanical compression (e.g. for severe epistaxis), surgical haemostasis with bleeding control procedures, fluid replacement and haemodynamic support, blood products (packed red cells or fresh frozen plasma, depending on associated anaemia or coagulopathy) or platelets.

For life-threatening bleeding that cannot be controlled with the measures such as transfusion or haemostasis, the administration of a 4-factor prothrombin complex concentrate (PCC) at 50 IU/kg has been shown to reverse the effects of Lixiana 30 minutes after completing the infusion.

Recombinant factor VIIa (r-FVIIa) can also be considered. However, there is limited clinical experience with the use of this product in individuals receiving edoxaban.

Depending on local availability, a consultation with a coagulation expert should be considered in case of major bleedings.

Protamine sulfate and vitamin K are not expected to affect the anticoagulant activity of edoxaban.
There is no experience with antifibrinolytic agents (tranexamic acid, aminocaproic acid) in individuals receiving edoxaban. There is neither scientific rationale for benefit nor experience with the use of systemic haemostatics (desmopressin, aprotinin) in individuals receiving edoxaban. Due to the high plasma protein binding edoxaban is not expected to be dialysable.

5. **PHARMACOLOGICAL PROPERTIES**

5.1 **Pharmacodynamic properties**

Pharmacotherapeutic group: Other antithrombotic agents, ATC code: B01AF03

**Mechanism of action**

Edoxaban is a highly selective, direct and reversible inhibitor of factor Xa, the serine protease located in the final common pathway of the coagulation cascade. Edoxaban inhibits free factor Xa, and prothrombinase activity. Inhibition of factor Xa in the coagulation cascade reduces thrombin generation, prolongs clotting time and reduces the risk of thrombus formation.

**Pharmacodynamic effects**

Edoxaban produces rapid onset of pharmacodynamic effects within 1 - 2 hours, which corresponds with peak edoxaban exposure (Cmax). The pharmacodynamic effects measured by anti-factor Xa assay are predictable and correlate with the dose and the concentration of edoxaban. As a result of FXa inhibition, edoxaban also prolongs clotting time in tests such as prothrombin time (PT), and activated partial thromboplastin time (aPTT). Changes observed in these clotting tests are expected at the therapeutic dose, however, these changes are small, subject to a high degree of variability, and not useful in monitoring the anticoagulation effect of edoxaban.

**Effects of coagulation markers when switching from rivaroxaban, dabigatran, or apixaban to edoxaban**

In clinical pharmacology studies, healthy subjects received rivaroxaban 20 mg once daily, dabigatran 150 mg twice daily, or apixaban 5 mg twice daily, followed by a single dose of edoxaban 60 mg on Day 4. The effect on prothrombin time (PT) and other coagulation biomarkers (e.g. anti-FXa, aPTT) was measured. Following the switch to edoxaban on Day 4 the PT was equivalent to Day 3 of rivaroxaban and apixaban. For dabigatran higher aPTT activity was observed after edoxaban administration with prior dabigatran treatment compared to that after treatment with edoxaban alone. This is considered to be due to the carry-over effect of dabigatran treatment, however, this did not lead to a prolongation of bleeding time.

Based on these data, when switching from these anticoagulants to edoxaban, the first dose of edoxaban can be initiated at the time of the next scheduled dose of the previous anticoagulant (see section 4.2).

**Clinical efficacy and safety**

*Prevention of stroke and systemic embolism*

The edoxaban clinical programme for atrial fibrillation was designed to demonstrate the efficacy and safety of two dose groups of edoxaban compared to warfarin for the prevention of stroke and systemic embolism in subjects with nonvalvular atrial fibrillation and at moderate to high risk of stroke and systemic embolic events (SEE).

In the pivotal ENGAGE AF-TIMI 48 study (an event-driven, Phase 3, multi-centre, randomised, double-blind double-dummy parallel-group study), 21,105 subjects, with a mean CHADS2 score of 2.8, were randomised to either edoxaban 30 mg once daily treatment group, or edoxaban 60 mg once daily treatment group or warfarin. Subjects in both edoxaban treatment groups had their dose halved if one or more of the following clinical factors were present: moderate renal impairment (CrCL 30 – 50 mL/min), low body weight (≤ 60 kg) or concomitant use of specific P-gp inhibitors (verapamil, quinidine, dronedarone).
The primary efficacy endpoint was the composite of stroke and SEE. Secondary efficacy endpoints included: Composite of stroke, SEE, and cardiovascular (CV) mortality; major adverse cardiovascular event (MACE), which is the composite of non-fatal MI, non-fatal stroke, non-fatal SEE, and death due to CV cause or bleeding; composite of stroke, SEE, and all-cause mortality.

The median study drug exposure for both the edoxaban 60 mg and 30 mg treatment groups was 2.5 years. The median study follow-up for both the edoxaban 60 mg and 30 mg treatment groups was 2.8 years. The median subject-year exposure was 15,471, and 15,840 for the 60 mg and 30 mg treatment groups, respectively; and the median subject-year follow-up was 19,191 and 19,216 for the 60 mg and 30 mg treatment groups, respectively.

In the warfarin group, the median TTR (time in therapeutic range, INR 2.0 to 3.0) was 68.4%.

The main analysis of efficacy was aimed to show the non-inferiority of edoxaban versus warfarin on first stroke or SEE that occurred during treatment or within 3 days from the last dose taken in the modified intention-to-treat (mITT) population. Edoxaban 60 mg was non-inferior to warfarin for the primary efficacy endpoint of stroke or SEE (upper limit of the 97.5% CI of the HR was below the pre-specified non-inferiority margin of 1.38) (Table 4).

<table>
<thead>
<tr>
<th>Primary Endpoint</th>
<th>Edoxaban 60 mg (30 mg Dose Reduced) (N = 7,012)</th>
<th>Warfarin (N = 7,012)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>First Stroke/SEE</strong>&lt;sup&gt;a&lt;/sup&gt;</td>
<td></td>
<td></td>
</tr>
<tr>
<td>n</td>
<td>182</td>
<td>232</td>
</tr>
<tr>
<td>Event Rate (%/yr)&lt;sup&gt;b&lt;/sup&gt;</td>
<td>1.18</td>
<td>1.50</td>
</tr>
<tr>
<td>HR (97.5% CI)</td>
<td>0.79 (0.63, 0.99)</td>
<td></td>
</tr>
<tr>
<td>p-value for non-inferiority&lt;sup&gt;c&lt;/sup&gt;</td>
<td>&lt;0.0001</td>
<td></td>
</tr>
<tr>
<td><strong>First Ischaemic Stroke</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>n</td>
<td>135</td>
<td>144</td>
</tr>
<tr>
<td>Event Rate (%/yr)&lt;sup&gt;b&lt;/sup&gt;</td>
<td>0.87</td>
<td>0.93</td>
</tr>
<tr>
<td>HR (95% CI)</td>
<td>0.94 (0.75, 1.19)</td>
<td></td>
</tr>
<tr>
<td><strong>First Haemorrhagic Stroke</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>n</td>
<td>40</td>
<td>76</td>
</tr>
<tr>
<td>Event Rate (%/yr)&lt;sup&gt;b&lt;/sup&gt;</td>
<td>0.26</td>
<td>0.49</td>
</tr>
<tr>
<td>HR (95% CI)</td>
<td>0.53 (0.36, 0.78)</td>
<td></td>
</tr>
<tr>
<td><strong>First SEE</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>n (%/yr)&lt;sup&gt;b&lt;/sup&gt;</td>
<td>8 (0.05)</td>
<td>13 (0.08)</td>
</tr>
<tr>
<td>HR (95% CI)</td>
<td>0.62 (0.26, 1.50)</td>
<td></td>
</tr>
</tbody>
</table>

Abbreviations: HR = Hazard Ratio versus warfarin, CI = Confidence Interval, n = number of events, mITT = modified Intent To Treat, N = number of subjects in mITT population, SEE = Systemic Embolic Event, yr = year.

<sup>a</sup> A subject can be represented in multiple rows.

<sup>b</sup> The event rate (%/yr) is calculated as number of events/subject-year exposure.

<sup>c</sup> The two-sided p-value is based on the non-inferiority margin of 1.38.

During the overall study period in the ITT population (analysis set to show superiority), adjudicated stroke or SEE occurred in 296 subjects in the edoxaban 60 mg group (1.57% per year), and 337
subjects in the warfarin group (1.80% per year). Compared to warfarin-treated subjects, the HR in the edoxaban 60 mg group was 0.87 (99% CI: 0.71, 1.07, p = 0.08 for superiority).

In subgroup analyses, for subjects in the 60 mg treatment group who were dose reduced to 30 mg in the ENGAGE AF-TIMI 48 study (for body weight \( \leq 60 \) kg, moderate renal impairment, or concomitant use of P-gp inhibitors), the event rate was: 2.29% per year for the primary endpoint, compared to the event rate of 2.66% per year for the matching subjects in the warfarin group [HR (95% CI): 0.86 (0.66, 1.13)].

The efficacy results for pre-specified major subgroups (with dose reduction as required), including age, body weight, gender, status of renal function, prior stroke or TIA, diabetes and P-gp inhibitors were generally consistent with the primary efficacy results for the overall population studied in the trial.

The Hazard Ratio (Edoxaban 60 mg vs. warfarin) for the primary endpoint in the centres with a lower average time of INR in the target range (INR TTR) for warfarin was 0.73 – 0.80 for the lowest 3 quartiles (INR TTR \( \leq 57.7\% \) to \( \leq 73.9\% \)). It was 1.07 in centres with the best control of warfarin therapy (4th quartile with \( > 73.9\% \) of INR values in the therapeutic range).

There was a statistically significant interaction between the effect of edoxaban versus warfarin on the main study outcome (stroke/SEE) and renal function (p-value 0.0042; mITT, overall study period).

Table 5 shows ischaemic strokes/SEE by creatinine clearance category in NVAF patients in ENGAGE AF-TIMI 48. There is a decreasing event rate at increasing CrCL in both treatment groups.

Table 5: Number of Ischaemic Strokes/SEE by creatinine clearance category in ENGAGE AF-TIMI 48, mITT Analysis Set Overall Study

<table>
<thead>
<tr>
<th>CrCL subgroup (mL/min)</th>
<th>Edoxaban 60 mg (N = 7,012)</th>
<th>Warfarin (N = 7,012)</th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>n</td>
<td>Number of Events</td>
<td>Event rate (%/year)</td>
<td>n</td>
<td>Number of Events</td>
</tr>
<tr>
<td>≥ 30 to ≤ 50</td>
<td>1,302</td>
<td>63</td>
<td>1.89</td>
<td>1,305</td>
<td>67</td>
</tr>
<tr>
<td>&gt; 50 to ≤ 70</td>
<td>2,093</td>
<td>85</td>
<td>1.51</td>
<td>2,106</td>
<td>95</td>
</tr>
<tr>
<td>&gt; 70 to ≤ 90</td>
<td>1,661</td>
<td>45</td>
<td>0.99</td>
<td>1,703</td>
<td>50</td>
</tr>
<tr>
<td>&gt; 90 to ≤ 110</td>
<td>927</td>
<td>27</td>
<td>1.08</td>
<td>960</td>
<td>26</td>
</tr>
<tr>
<td>&gt; 110 to ≤ 130</td>
<td>497</td>
<td>14</td>
<td>1.01</td>
<td>469</td>
<td>10</td>
</tr>
<tr>
<td>&gt; 130</td>
<td>462</td>
<td>10</td>
<td>0.78</td>
<td>418</td>
<td>3</td>
</tr>
</tbody>
</table>

Abbreviations: N = number of subjects; mITT population overall study period; n = number of patients in subgroup
*HR not computed if number of events < 5 in one treatment group.

Within renal function subgroups, results for the secondary efficacy endpoints were consistent with those for the primary endpoint.

Superiority testing was performed on the ITT Overall Study Period. Stroke and SEE occurred in fewer subjects in the edoxaban 60 mg treatment group than in the warfarin group (1.57% and 1.80% per year, respectively), with a HR of 0.87 (99% CI: 0.71, 1.07, p = 0.0807 for superiority).
The pre-specified composite endpoints for the comparison of the edoxaban 60 mg treatment group to warfarin for stroke, SEE, and CV mortality HR (99% CI) was 0.87 (0.76, 0.99), MACE 0.89 (0.78, 1.00), and stroke, SEE, and all-cause mortality 0.90 (0.80, 1.01).

The results for all-cause mortality (adjudicated deaths) in the ENGAGE AF-TIMI 48 study were 769 (3.99% per year) for subjects taking edoxaban 60 mg (30 mg dose reduced) as opposed to 836 (4.35% per year) for warfarin [HR (95% CI): 0.91 (0.83, 1.01)]. All-cause mortality (adjudicated deaths) per renal subgroups (edoxaban vs. warfarin): CrCL 30 to ≤ 50 mL/min [HR (95% CI): 0.81 (0.68, 0.97)]; CrCL > 50 to < 80 mL/min [HR (95% CI): 0.87 (0.75, 1.02)]; CrCL ≥ 80 mL/min [HR (95% CI): 1.15 (0.95, 1.40)].

Edoxaban 60 mg (30 mg dose reduced) resulted in a lower rate of cardiovascular mortality compared to warfarin [HR (95% CI): 0.86 (0.77, 0.97)]. Adjudicated efficacy cardiovascular mortality per renal subgroups (edoxaban vs. warfarin): CrCL 30 to ≤ 50 mL/min [HR (95% CI): 0.80 (0.65, 0.99)]; CrCL > 50 to < 80 mL/min [HR (95% CI): 0.75 (0.62, 0.90)]; CrCL ≥ 80 mL/min [HR (95% CI): 1.16 (0.92, 1.46)].

Safety in patients with NVAF in ENGAGE AF-TIMI 48

The primary safety endpoint was major bleeding.

There was a significant risk reduction in favour of the edoxaban 60 mg treatment group compared with the warfarin group in major bleeding (2.75%, and 3.43% per year, respectively) [HR (95% CI): 0.80 (0.71, 0.91); p = 0.0009], ICH (0.39%, and 0.85% per year, respectively) [HR (95% CI): 0.47 (0.34, 0.63); p < 0.0001], and other types of bleeding (Table 6).

The reduction in fatal bleeds was also significant for the edoxaban 60 mg treatment group compared with the warfarin group (0.21%, and 0.38%) [HR (95% CI): 0.55 (0.36, 0.84); p = 0.0059 for superiority], primarily because of the reduction in fatal ICH bleeds [HR (95% CI): 0.58 (0.35, 0.95); p = 0.0312].
Table 6: Bleeding Events in ENGAGE AF-TIMI 48 Study - Safety Analysis On-Treatment

<table>
<thead>
<tr>
<th></th>
<th>Edoxaban 60 mg (30 mg Dose Reduced) (N = 7,012)</th>
<th>Warfarin (N = 7,012)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Major Bleeding</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>n</td>
<td>418</td>
<td>524</td>
</tr>
<tr>
<td>Event rate (%/yr)(^a)</td>
<td>2.75</td>
<td>3.43</td>
</tr>
<tr>
<td>HR (95% CI)</td>
<td>0.80 (0.71, 0.91)</td>
<td></td>
</tr>
<tr>
<td>p-value</td>
<td>0.0009</td>
<td></td>
</tr>
<tr>
<td>ICH(^b)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>n</td>
<td>61</td>
<td>132</td>
</tr>
<tr>
<td>Event rate (%/yr)(^a)</td>
<td>0.39</td>
<td>0.85</td>
</tr>
<tr>
<td>HR (95% CI)</td>
<td>0.47 (0.34, 0.63)</td>
<td></td>
</tr>
<tr>
<td><strong>Fatal Bleeding</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>n</td>
<td>32</td>
<td>59</td>
</tr>
<tr>
<td>Event rate (%/yr)(^a)</td>
<td>0.21</td>
<td>0.38</td>
</tr>
<tr>
<td>HR (95% CI)</td>
<td>0.55 (0.36, 0.84)</td>
<td></td>
</tr>
<tr>
<td><strong>CRNM Bleeding</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>n</td>
<td>1,214</td>
<td>1,396</td>
</tr>
<tr>
<td>Event rate (%/yr)(^a)</td>
<td>8.67</td>
<td>10.15</td>
</tr>
<tr>
<td>HR (95% CI)</td>
<td>0.86 (0.80, 0.93)</td>
<td></td>
</tr>
<tr>
<td><strong>Any Confirmed Bleeding(^c)</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>n</td>
<td>1,865</td>
<td>2,114</td>
</tr>
<tr>
<td>Event rate (%/yr)(^a)</td>
<td>14.15</td>
<td>16.40</td>
</tr>
<tr>
<td>HR (95% CI)</td>
<td>0.87 (0.82, 0.92)</td>
<td></td>
</tr>
</tbody>
</table>

Abbreviations: ICH = Intracranial Haemorrhage, HR = Hazard Ratio versus warfarin, CI = Confidence Interval, CRNM = Clinically Relevant Non-Major, n = number of subjects with events, N = number of subjects in Safety population, yr = year.

\(^a\) The event rate (%/yr) is calculated as number of events/subject-year exposure.

\(^b\) ICH includes primary haemorrhagic stroke, subarachnoid haemorrhage, epi-/subdural haemorrhage, and ischaemic stroke with major haemorrhagic conversion. All ICHs reported on the Adjudicated Cerebrovascular and Non-Intracranial bleed eCRF forms confirmed by the adjudicators are included in ICH counts.

\(^c\) 'Any Confirmed Bleeding' includes those that the adjudicator defined as clinically overt.

Note: A subject can be included in multiple sub-categories if he/she had an event for those categories. The first event of each category is included in the analysis.

Tables 7, 8 and 9 show major, fatal and intracranial bleedings, respectively, by creatinine clearance category in NVAF patients in ENGAGE AF-TIMI 48. There is a decreasing event rate at increasing CrCL in both treatment groups.
### Table 7: Number of Major Bleeding Events by creatinine clearance category in ENGAGE AF-TIMI 48, Safety Analysis On-Treatment<sup>a</sup>

<table>
<thead>
<tr>
<th>CrCL subgroup (mL/min)</th>
<th>Edoxaban 60 mg (N = 7,012)</th>
<th>Warfarin (N = 7,012)</th>
<th>HR (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>n</td>
<td>Number of Events</td>
<td>Event rate (%/year)</td>
</tr>
<tr>
<td>≥ 30 to ≤ 50</td>
<td>1,302</td>
<td>96</td>
<td>3.91</td>
</tr>
<tr>
<td>&gt; 50 to ≤ 70</td>
<td>2,093</td>
<td>148</td>
<td>3.31</td>
</tr>
<tr>
<td>&gt; 70 to ≤ 90</td>
<td>1,661</td>
<td>108</td>
<td>2.88</td>
</tr>
<tr>
<td>&gt; 90 to ≤ 110</td>
<td>927</td>
<td>29</td>
<td>1.33</td>
</tr>
<tr>
<td>&gt; 110 to ≤ 130</td>
<td>497</td>
<td>20</td>
<td>1.70</td>
</tr>
<tr>
<td>&gt; 130</td>
<td>462</td>
<td>13</td>
<td>1.18</td>
</tr>
</tbody>
</table>

### Table 8: Number of Fatal Bleeding Events by creatinine clearance category in ENGAGE AF-TIMI 48, Safety Analysis On-Treatment<sup>a</sup>

<table>
<thead>
<tr>
<th>CrCL subgroup (mL/min)</th>
<th>Edoxaban 60 mg (N = 7,012)</th>
<th>Warfarin (N = 7,012)</th>
<th>HR (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>n</td>
<td>Number of Events</td>
<td>Event rate (%/year)</td>
</tr>
<tr>
<td>≥ 30 to ≤ 50</td>
<td>1,302</td>
<td>9</td>
<td>0.36</td>
</tr>
<tr>
<td>&gt; 50 to ≤ 70</td>
<td>2,093</td>
<td>8</td>
<td>0.18</td>
</tr>
<tr>
<td>&gt; 70 to ≤ 90</td>
<td>1,661</td>
<td>10</td>
<td>0.26</td>
</tr>
<tr>
<td>&gt; 90 to ≤ 110</td>
<td>927</td>
<td>2</td>
<td>0.09</td>
</tr>
<tr>
<td>&gt; 110 to ≤ 130</td>
<td>497</td>
<td>1</td>
<td>0.08</td>
</tr>
<tr>
<td>&gt; 130</td>
<td>462</td>
<td>2</td>
<td>0.18</td>
</tr>
</tbody>
</table>

### Table 9: Number of Intracranial Bleeding Events by creatinine clearance category in ENGAGE AF-TIMI 48, Safety Analysis On-Treatment<sup>a</sup>

<table>
<thead>
<tr>
<th>CrCL subgroup (mL/min)</th>
<th>Edoxaban 60 mg (N = 7,012)</th>
<th>Warfarin (N = 7,012)</th>
<th>HR (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>n</td>
<td>Number of Events</td>
<td>Event rate (%/year)</td>
</tr>
<tr>
<td>≥ 30 to ≤ 50</td>
<td>1,302</td>
<td>16</td>
<td>0.64</td>
</tr>
<tr>
<td>&gt; 50 to ≤ 70</td>
<td>2,093</td>
<td>19</td>
<td>0.42</td>
</tr>
<tr>
<td>&gt; 70 to ≤ 90</td>
<td>1,661</td>
<td>17</td>
<td>0.44</td>
</tr>
<tr>
<td>&gt; 90 to ≤ 110</td>
<td>927</td>
<td>5</td>
<td>0.23</td>
</tr>
<tr>
<td>&gt; 110 to ≤ 130</td>
<td>497</td>
<td>2</td>
<td>0.17</td>
</tr>
<tr>
<td>&gt; 130</td>
<td>462</td>
<td>1</td>
<td>0.09</td>
</tr>
</tbody>
</table>

Abbreviations: N = number of subjects; mITT population overall study period; n = number of patients in subgroup

*HR not computed if number of events < 5 in one treatment group.

<sup>a</sup> On-Treatment: Time from first dose of study drug to last dose plus 3 days.
In subgroup analyses, for subjects in the 60 mg treatment group who were dose reduced to 30 mg in the ENGAGE AF-TIMI 48 study for body weight ≤ 60 kg, moderate renal impairment, or concomitant use of P-gp inhibitors, 104 (3.05% per year) of edoxaban 30 mg dose reduced subjects and 166 (4.85% per year) of warfarin dose reduced subjects had a major bleeding event [HR (95% CI): 0.63 (0.50, 0.81)].

In the ENGAGE AF-TIMI 48 study there was a significant improvement in Net Clinical Outcome (First Stroke, SEE, Major Bleed, or All-Cause Mortality; mITT population, overall study period) in favour of edoxaban, HR (95% CI): 0.89 (0.83, 0.96); p = 0.0024, when edoxaban 60 mg treatment group was compared to warfarin.

Treatment of DVT, treatment of PE and the prevention of recurrent DVT and PE (VTE)
The edoxaban clinical programme for VTE was designed to demonstrate the efficacy and safety of edoxaban in the treatment of DVT and PE, and the prevention of recurrent DVT and PE.

In the pivotal Hokusai-VTE study, 8,292 subjects were randomised to receive initial heparin therapy (enoxaparin or unfractionated heparin) followed by edoxaban 60 mg once daily or the comparator. In the comparator arm, subjects received initial heparin therapy concurrently with warfarin, titrated to a target INR of 2.0 to 3.0, followed by warfarin alone. The treatment duration was from 3 months up to 12 months, determined by the investigator based on the patient’s clinical features.

The majority of edoxaban treated patients were Caucasians (69.6%) and Asians (21.0%), 3.8% were Black, 5.3% were categorised as Other race.

The duration of therapy was at least 3 months for 3,718 (91.6%) edoxaban subjects versus 3,727 (91.4%) of warfarin subjects; at least 6 months for 3,495 (86.1%) of edoxaban subjects versus 3,491 (85.6%) of warfarin subjects; and 12 months for 1,643 (40.5%) edoxaban subjects versus 1,659 (40.4%) of warfarin subjects.

The primary efficacy endpoint was the recurrence of symptomatic VTE, defined as the composite of recurrent symptomatic DVT, non-fatal symptomatic PE and fatal PE in subjects during the 12-month study period. Secondary efficacy outcomes included the composite clinical outcome of recurrent VTE and all-cause mortality.

Edoxaban 30 mg once daily was used for subjects with one or more of the following clinical factors: moderate renal impairment (CrCL 30 - 50 mL/min); body weight ≤ 60 kg; concomitant use of specific P-gp inhibitors.

In the Hokusai-VTE study (Table 10) edoxaban was demonstrated to be non-inferior to warfarin for the primary efficacy outcome, recurrent VTE, which occurred in 130 of 4,118 subjects (3.2%) in the edoxaban group versus 146 of 4,122 subjects (3.5%) in the warfarin group [HR (95% CI): 0.89 (0.70, 1.13); p < 0.0001 for non-inferiority]. In the warfarin group, the median TTR (time in therapeutic range, INR 2.0 to 3.0) was 65.6%. For subjects presenting with PE (with or without DVT), 47 (2.8%) of edoxaban and 65 (3.9%) of warfarin subjects had a recurrent VTE [HR (95% CI): 0.73 (0.50, 1.06)].
Table 10: Efficacy Results from the Hokusai-VTE Study - mITT population, overall study period

<table>
<thead>
<tr>
<th>Primary endpointa</th>
<th>Edoxaban 60 mg (30 mg Dose Reduced) (N = 4,118)</th>
<th>Warfarin (N = 4,122)</th>
<th>Edoxaban vs Warfarin HR (95% CI)b p-valuec</th>
</tr>
</thead>
<tbody>
<tr>
<td>All subjects with symptomatic recurrent VTE, n (%)</td>
<td>130 (3.2)</td>
<td>146 (3.5)</td>
<td>0.89 (0.70, 1.13) p-value &lt; 0.0001 (non-inferiority)</td>
</tr>
<tr>
<td>PE with or without DVT</td>
<td>73 (1.8)</td>
<td>83 (2.0)</td>
<td></td>
</tr>
<tr>
<td>Fatal PE or Death where PE cannot be ruled out</td>
<td>24 (0.6)</td>
<td>24 (0.6)</td>
<td></td>
</tr>
<tr>
<td>Non-fatal PE</td>
<td>49 (1.2)</td>
<td>59 (1.4)</td>
<td></td>
</tr>
<tr>
<td>DVT only</td>
<td>57 (1.4)</td>
<td>63 (1.5)</td>
<td></td>
</tr>
</tbody>
</table>

Abbreviations: CI = Confidence Interval; DVT = deep vein thrombosis; mITT = modified intent-to-treat; HR = Hazard Ratio vs. warfarin; n = number of subjects with events; N = number of subjects in mITT population; PE = pulmonary embolism; VTE = venous thromboembolic events.

a The primary efficacy endpoint is adjudicated symptomatic recurrent VTE (i.e., the composite endpoint of DVT, non-fatal PE, and fatal PE).
b The HR, two-sided CI are based on the Cox proportional hazards regression model including treatment and the following randomisation stratification factors as covariates: presenting diagnosis (PE with or without DVT, DVT only), baseline risk factors (temporary factors, all others), and the need for 30 mg edoxaban/edoxaban placebo dose at randomisation (yes/no).
c The p-value is for the pre-defined non-inferiority margin of 1.5.

For the subjects who were dose reduced to 30 mg (predominantly low body weight or renal function) 15 (2.1%) edoxaban and 22 (3.1%) of warfarin subjects had a recurrent VTE [HR (95% CI): 0.69 (0.36, 1.34)].

The secondary composite endpoint of recurrent VTE and all-cause mortality occurred in 138 subjects (3.4%) in the edoxaban group and 158 subjects (3.9%) in the warfarin group [HR (95% CI): 0.87 (0.70, 1.10)].

The results for all-cause mortality (adjudicated deaths) in Hokusai-VTE were 136 (3.3%) for subjects taking edoxaban 60 mg (30 mg dose reduced) as opposed to 130 (3.2%) for warfarin.

In a pre-specified subgroup analysis of PE subjects 447 (30.6%) and 483 (32.2%) of edoxaban and warfarin treated subjects, respectively, were identified as having PE and NT-proBNP ≥ 500 pg/mL. The primary efficacy outcome occurred in 14 (3.1%) and 30 (6.2%) of edoxaban and warfarin subjects, respectively [HR (95% CI): 0.50 (0.26, 0.94)].

The efficacy results for pre-specified major subgroups (with dose reduction as required), including age, body weight, gender and status of renal function were consistent with the primary efficacy results for the overall population studied in the trial.

Safety in patients with VTE (DVT and PE) in Hokusai-VTE

The primary safety endpoint was clinically relevant bleeding (major or clinically relevant non-major).

Table 11 summarises adjudicated bleeding events for the safety analysis set on-treatment period.
There was a significant risk reduction in favour of edoxaban compared with warfarin for the primary safety endpoint of clinically relevant bleeding, a composite of major bleeding or clinically relevant non-major bleeding (CRNM), which occurred in 349 of 4,118 subjects (8.5%) in the edoxaban group and in 423 of 4,122 subjects (10.3%) in the warfarin group [HR (95% CI): 0.81 (0.71, 0.94); p = 0.004 for superiority].

Table 11: Bleeding Events in Hokusai-VTE Study - Safety Analysis On-Treatment Perioda

<table>
<thead>
<tr>
<th></th>
<th>Edoxaban 60 mg (30 mg Dose Reduced) (N = 4,118)</th>
<th>Warfarin (N = 4,122)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Clinically Relevant Bleeding (Major and CRNM)b, n (%)</td>
<td>349 (8.5)</td>
<td>423 (10.3)</td>
</tr>
<tr>
<td>HR (95% CI)</td>
<td>0.81 (0.71, 0.94)</td>
<td></td>
</tr>
<tr>
<td>p-value</td>
<td>0.004 (for superiority)</td>
<td></td>
</tr>
<tr>
<td>Major Bleeding n (%)</td>
<td>56 (1.4)</td>
<td>66 (1.6)</td>
</tr>
<tr>
<td>HR (95% CI)</td>
<td>0.84 (0.59, 1.21)</td>
<td></td>
</tr>
<tr>
<td>ICH fatal</td>
<td>0</td>
<td>6 (0.1)</td>
</tr>
<tr>
<td>ICH non-fatal</td>
<td>5 (0.1)</td>
<td>12 (0.3)</td>
</tr>
<tr>
<td>CRNM Bleeding n (%)</td>
<td>298 (7.2)</td>
<td>368 (8.9)</td>
</tr>
<tr>
<td>HR (95% CI)</td>
<td>0.80 (0.68, 0.93)</td>
<td></td>
</tr>
<tr>
<td>All Bleeding n (%)</td>
<td>895 (21.7)</td>
<td>1,056 (25.6)</td>
</tr>
<tr>
<td>HR (95% CI)</td>
<td>0.82 (0.75, 0.90)</td>
<td></td>
</tr>
</tbody>
</table>

Abbreviations: ICH = Intracranial Haemorrhage, HR = Hazard Ratio vs. warfarin; CI = Confidence Interval; N = number of subjects in safety population; n = number of events; CRNM = clinically relevant non-major

a On-Treatment Period: Time from first dose of study drug to last dose plus 3 days.
b Primary Safety Endpoint: Clinically relevant bleeding (composite of major and clinically relevant non-major bleeding).

In subgroup analyses, for subjects who were dose reduced to 30 mg in the Hokusai-VTE study for body weight ≤ 60 kg, moderate renal impairment, or concomitant use of P-gp inhibitors, 58 (7.9%) of edoxaban 30 mg dose reduced subjects and 92 (12.8%) of warfarin subjects had a major bleeding or CRNM event [HR (95%): 0.62 (0.44, 0.86)].

In the Hokusai-VTE study the Net Clinical Outcome (Recurrent VTE, Major Bleed, or All-Cause Mortality; mITT population, overall study period) HR (95% CI) was 1.00 (0.85, 1.18) when edoxaban was compared to warfarin.

Patients undergoing cardioversion
A multicentre, prospective, randomised, open-label study with blinded endpoint evaluation (ENSURE-AF) was conducted which randomised 2199 subjects (oral anticoagulant naïve and pre-treated) with non-valvular atrial fibrillation scheduled for cardioversion, to compare edoxaban 60 mg once daily with enoxaparin/warfarin to maintain a therapeutic INR of 2.0 to 3.0 (randomised 1:1), mean TTR on warfarin was 70.8%. A total of 2149 subjects were treated with either edoxaban (N = 1067) or enoxaparin/warfarin (N = 1082). Subjects in the edoxaban treatment group received 30 mg once daily
if one or more of the following clinical factors were present: moderate renal impairment (CrCL 30 – 50 mL/min), low body weight (≤ 60 kg) or concomitant use of specific P-gp inhibitors. The majority of subjects in the edoxaban and warfarin groups had cardioversion performed (83.7% and 78.9%, respectively) or were auto-converted (6.6% and 8.6%, respectively). TEE-guided (within 3 days of initiation) or conventional cardioversion (at least 21 days of pre-treatment) was employed. Subjects were maintained on treatment for 28 days post cardioversion.

The primary efficacy outcome consisted of a composite of all stroke, SEE, MI and CV mortality. A total of 5 (0.5%, 95% CI 0.15% - 1.06%) events occurred in subjects in the edoxaban group (N = 1095) and 11 (1.0%, 95% CI 0.50% - 1.78%) events in the warfarin group (N = 1104); OR 0.46 (95% CI 0.12 - 1.43); ITT analysis set overall study period with mean duration of 66 days.

The primary safety outcome was a composite of major and CRNM bleeding. A total of 16 (1.5%, 95% CI 0.86% - 2.42%) events occurred in subjects in the edoxaban (N = 1067) group and 11 (1.0%, 95% CI 0.51% - 1.81%) events in the warfarin (N = 1082) group; OR 1.48 (95% CI 0.64 - 3.55); safety analysis set on-treatment period.

This exploratory study showed low rates of major and CRNM bleeding and thromboembolism in the two treatment groups in the setting of cardioversion.

**Paediatric population**

The European Medicines Agency has deferred the obligation to submit the results of studies with edoxaban in one or more subsets of the paediatric population in prevention of arterial thrombosis, treatment of thromboembolism and prevention of thromboembolism (see section 4.2 for information on paediatric use).

### 5.2 Pharmacokinetic properties

**Absorption**

Edoxaban is absorbed with peak plasma concentrations within 1 - 2 hours. The absolute bioavailability is approximately 62%. Food increases peak exposure to a varying extent, but has minimal effect on total exposure. Edoxaban was administered with or without food in the ENGAGE AF-TIMI 48 and the Hokusai-VTE studies. Edoxaban is poorly soluble at pH of 6.0 or higher. Co-administration of proton-pump inhibitors had no relevant impact on edoxaban exposure.

**Distribution**

Disposition is biphasic. The volume of distribution is 107 (19.9) L mean (SD). In vitro plasma protein binding is approximately 55%. There is no clinically relevant accumulation of edoxaban (accumulation ratio 1.14) with once daily dosing. Steady state concentrations are achieved within 3 days.

**Biotransformation**

Unchanged edoxaban is the predominant form in plasma. Edoxaban is metabolised via hydrolysis (mediated by carboxylesterase 1), conjugation or oxidation by CYP3A4/5 (< 10%). Edoxaban has three active metabolites, the predominant metabolite (M-4), formed by hydrolysis, is active and reaches less than 10% of the exposure of the parent compound in healthy subjects. Exposure to the other metabolites is less than 5%. Edoxaban is a substrate for the efflux transporter P-glycoprotein (P-gp), but not a substrate for uptake transporters such as organic anion transporter polypeptide OATP1B1, organic anion transporters OAT1 or OAT3 or organic cation transporter OCT2. Its active metabolite is a substrate for OATP1B1.

**Elimination**

In healthy subjects, the total clearance is estimated as 22 (± 3) L/hour; 50% is renally cleared (11 L/hour). Renal clearance accounts for approximately 35% of the administered dose. Metabolism and biliary/intestinal excretion account for the remaining clearance. The t½ for oral administration is 10 - 14 hours.
**Linearity/non-linearity**

Edoxaban displays approximately dose-proportional pharmacokinetics for doses of 15 mg to 60 mg in healthy subjects.

**Special populations**

**Elderly**

After taking renal function and body weight into account, age had no additional clinically significant effect on edoxaban pharmacokinetics in a population pharmacokinetic analysis of the pivotal Phase 3 study in NVAF (ENGAGE AF-TIMI 48).

**Gender**

After accounting for body weight, gender had no additional clinically significant effect on edoxaban pharmacokinetics in a population pharmacokinetic analysis of the Phase 3 study in NVAF (ENGAGE AF-TIMI 48).

**Ethnic origin**

In a population pharmacokinetic analysis of the ENGAGE AF-TIMI 48 study, peak and total exposure in Asian patients and non-Asian patients were comparable.

**Renal impairment**

The plasma AUCs for subjects with mild (CrCL > 50 - 80 mL/min), moderate (CrCL 30 - 50 mL/min) and severe (CrCL < 30 mL/min but not undergoing dialysis) renal impairment were increased by 32%, 74%, and 72%, respectively, relative to subjects with normal renal function. In patients with renal impairment the metabolite profile changes and a higher quantity of active metabolites are formed. There is a linear correlation between edoxaban plasma concentration and anti-FXa activity regardless of renal function. Subjects with ESRD undergoing peritoneal dialysis had 93% higher total exposure compared with healthy subjects. Population PK modeling indicates that exposure approximately doubles in patients with severe renal impairment (CrCL 15 – 29 mL/min) relative to patients with normal renal function.

**Anti-FXa activity by CrCL category**

Table 12 below shows the edoxaban anti-Factor Xa activity by CrCL category for each indication.
Table 12: Edoxaban Anti-FXa activity by creatinine clearance

<table>
<thead>
<tr>
<th>Edoxaban Dose</th>
<th>CrCL (mL/min)</th>
<th>Edoxaban Anti-FXa activity post-dose (IU/mL)¹</th>
<th>Edoxaban Anti-FXa activity pre-dose (IU/mL)²</th>
</tr>
</thead>
<tbody>
<tr>
<td>30 mg QD</td>
<td>≥ 30 to ≤ 50</td>
<td>2.92 [0.33 – 5.88]</td>
<td>0.53 [0.11 – 2.06]</td>
</tr>
<tr>
<td>60 mg QD*</td>
<td>&gt; 50 to ≤ 70</td>
<td>4.52 [0.38 – 7.64]</td>
<td>0.83 [0.16 – 2.61]</td>
</tr>
<tr>
<td></td>
<td>&gt; 70 to ≤ 90</td>
<td>4.12 [0.19 – 7.55]</td>
<td>0.68 [0.05 – 2.33]</td>
</tr>
<tr>
<td></td>
<td>&gt; 90 to ≤ 110</td>
<td>3.82 [0.36 – 7.39]</td>
<td>0.60 [0.14 – 3.57]</td>
</tr>
<tr>
<td></td>
<td>&gt; 110 to ≤ 130</td>
<td>3.16 [0.28 – 6.71]</td>
<td>0.41 [0.15 – 1.51]</td>
</tr>
<tr>
<td></td>
<td>&gt; 130</td>
<td>2.76 [0.12 – 6.10]</td>
<td>0.45 [0.00 – 3.10]</td>
</tr>
</tbody>
</table>

| Treatment of DVT, treatment of PE and prevention of recurrent DVT and PE (VTE) |
|-------------------------------|-----------------|-----------------|-----------------|
| 30 mg QD                      | ≥ 30 to ≤ 50    | 2.21 [0.14 – 4.47] | 0.22 [0.00 – 1.09] |
| 60 mg QD*                     | > 50 to ≤ 70    | 3.42 [0.19 – 6.13] | 0.34 [0.00 – 3.10] |
|                               | > 70 to ≤ 90    | 2.97 [0.24 – 5.82] | 0.24 [0.00 – 1.77] |
|                               | > 90 to ≤ 110   | 2.82 [0.14 – 5.31] | 0.20 [0.00 – 2.52] |
|                               | > 110 to ≤ 130  | 2.64 [0.13 – 5.57] | 0.17 [0.00 – 1.86] |
|                               | > 130           | 2.39 [0.10 – 4.92] | 0.13 [0.00 – 2.43] |

*Dose reduction to 30 mg for low body weight ≤ 60 kg or specific concomitant P-glycoprotein (P-gp) inhibitors
¹ Post-dose is equivalent to C_{max} (post-dose samples were drawn 1 – 3 hours after edoxaban administration)
² Pre-dose is equivalent to C_{min}

Although treatment with edoxaban does not require routine monitoring, the effect on anticoagulation can be estimated by a calibrated quantitative anti-Factor Xa assay which may be useful in exceptional situations where knowledge of edoxaban exposure may help to inform clinical decisions, e.g. overdose and emergency surgery (see also section 4.4).

*Haemodialysis*
A 4 hour haemodialysis session reduced total edoxaban exposures by less than 9%.

*Hepatic impairment*
Patients with mild or moderate hepatic impairment exhibited comparable pharmacokinetics and pharmacodynamics to their matched healthy control group. Edoxaban has not been studied in patients with severe hepatic impairment (see section 4.2).
Body weight
In a population pharmacokinetic analysis of the ENGAGE AF-TIMI 48 study in NVAF, Cₘₐₓ and AUC in patients with median low body weight (55 kg) were increased by 40% and 13%, respectively, as compared with patients with median high body weight (84 kg). In Phase 3 clinical studies (both NVAF and VTE indications) patients with body weight ≤ 60 kg had a 50% edoxaban dose reduction and had similar efficacy and less bleeding when compared to warfarin.

Pharmacokinetic/pharmacodynamic relationship(s)
PT, INR, aPTT and Anti-factor Xa correlate linearly with edoxaban concentrations.

5.3 Preclinical safety data

Non-clinical data reveal no special hazard for humans based on conventional studies of safety pharmacology, repeated dose toxicity, genotoxicity, carcinogenic potential, or phototoxicity.

Reproductive toxicology
Edoxaban showed vaginal haemorrhage at higher doses in rats and rabbits but had no effects in the reproductive performance of parent rats.

In rats, no effects on male or female fertility were seen.

In animal reproduction studies, rabbits showed increased incidence of gallbladder variations at a dosage of 200 mg/kg which is approximately 65 times the maximum recommended human dose (MRHD) of 60 mg/day based on total body surface area in mg/m². Increased post-implantation pregnancy losses occurred in rats at 300 mg/kg/day (approximately 49 times the MRHD) and in rabbits at 200 mg/kg/day (approximately 65 times the MRHD) respectively.

Edoxaban was excreted in the breast milk of lactating rats.

Environmental Risk Assessment (ERA)
The active substance edoxaban tosilate is persistent in the environment (for instructions on disposal see section 6.6).

6. PHARMACEUTICAL PARTICULARS

6.1 List of excipients

Tablet core:
Mannitol (E421)
Pregelatinised starch
Crospovidone
Hydroxypropylcellulose
Magnesium stearate (E470b)

Film-coat:
Hypromellose (E464)
Macrogol 8000
Titanium dioxide (E171)
Talc
Carnauba wax
Iron oxide red (E172)

6.2 Incompatibilities

Not applicable.
6.3 Shelf life

5 years

6.4 Special precautions for storage

This medicinal product does not require any special storage conditions.

6.5 Nature and contents of container

PVC/Aluminium blisters. Cartons of 10, 14, 28, 30, 56, 60, 84, 90, 98, 100 film-coated tablets. PVC/Aluminium perforated unit dose blisters of 10 x 1, 50 x 1 and 100 x 1 film-coated tablets.

Not all pack sizes may be marketed.

6.6 Special precautions for disposal

Any unused medicinal product or waste material should be disposed of in accordance with local requirements.

7. MARKETING AUTHORISATION HOLDER

Daiichi Sankyo Europe GmbH
Zielstattstrasse 48
81379 Munich
Germany

8. MARKETING AUTHORISATION NUMBER(S)

EU/1/15/993/002
EU/1/15/993/004-015

9. DATE OF FIRST AUTHORISATION/RENEWAL OF THE AUTHORIZAITION

Date of first authorisation: 19 June 2015

10. DATE OF REVISION OF THE TEXT

Detailed information on this medicinal product is available on the website of the European Medicines Agency http://www.ema.europa.eu.
This medicinal product is subject to additional monitoring. This will allow quick identification of new safety information. Healthcare professionals are asked to report any suspected adverse reactions. See section 4.8 for how to report adverse reactions.

1. NAME OF THE MEDICINAL PRODUCT

Lixiana 60 mg film-coated tablets

2. QUALITATIVE AND QUANTITATIVE COMPOSITION

Each film-coated tablet contains 60 mg edoxaban (as tosilate).

For the full list of excipients, see section 6.1.

3. PHARMACEUTICAL FORM

Film-coated tablet.
Yellow, round-shaped film-coated tablets (10.5 mm diameter) debossed with “DSC L60”.

4. CLINICAL PARTICULARS

4.1 Therapeutic indications

Prevention of stroke and systemic embolism in adult patients with nonvalvular atrial fibrillation (NVAF) with one or more risk factors, such as congestive heart failure, hypertension, age ≥ 75 years, diabetes mellitus, prior stroke or transient ischaemic attack (TIA).

Treatment of deep vein thrombosis (DVT) and pulmonary embolism (PE), and prevention of recurrent DVT and PE in adults (see section 4.4 for haemodynamically unstable PE patients).

4.2 Posology and method of administration

Posology

Prevention of stroke and systemic embolism
The recommended dose is 60 mg edoxaban once daily.
Therapy with edoxaban in NVAF patients should be continued long term.

Treatment of DVT, treatment of PE and prevention of recurrent DVT and PE (VTE)
The recommended dose is 60 mg edoxaban once daily following initial use of parenteral anticoagulant for at least 5 days (see section 5.1). Edoxaban and initial parenteral anticoagulant should not be administered simultaneously.
The duration of therapy for treatment of DVT and PE (venous thromboembolism, VTE), and prevention of recurrent VTE should be individualised after careful assessment of the treatment benefit against the risk for bleeding (see section 4.4). Short duration of therapy (at least 3 months) should be based on transient risk factors (e.g. recent surgery, trauma, immobilisation) and longer durations should be based on permanent risk factors or idiopathic DVT or PE.
For NVAF and VTE the recommended dose is 30 mg edoxaban once daily in patients with one or more of the following clinical factors:

- Moderate or severe renal impairment (creatinine clearance (CrCL) 15 - 50 mL/min)
- Low body weight ≤ 60 kg
- Concomitant use of the following P-glycoprotein (P-gp) inhibitors: ciclosporin, dronedarone, erythromycin, or ketoconazole.

**Table 1: Summary of posology in NVAF and VTE (DVT and PE)**

<table>
<thead>
<tr>
<th>Summary Guide for Dosing</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Recommended dose</strong></td>
<td>60 mg once daily</td>
</tr>
<tr>
<td><strong>Dose recommendation for patients with one or more of the following clinical factors:</strong></td>
<td></td>
</tr>
<tr>
<td>Renal Impairment</td>
<td>Moderate or severe (CrCL 15 – 50 mL/min)</td>
</tr>
<tr>
<td>Low Body Weight</td>
<td>≤ 60 kg</td>
</tr>
<tr>
<td>P-gp Inhibitors</td>
<td>Ciclosporin, dronedarone, erythromycin, ketoconazole</td>
</tr>
</tbody>
</table>

**Missed dose**
If a dose of Lixiana is missed, the dose should be taken immediately and then be continued the following day with the once-daily intake as recommended. The patient should not take double the prescribed dose on the same day to make up for a missed dose.

**Switching to and from Lixiana**
Continued anticoagulant therapy is important in patients with NVAF and VTE. There may be situations that warrant a change in anticoagulation therapy (Table 2).

**Table 2: Switching**

<table>
<thead>
<tr>
<th>Switching to Lixiana</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>From</strong></td>
<td><strong>To</strong></td>
</tr>
<tr>
<td>Vitamin K Antagonist (VKA)</td>
<td>Lixiana</td>
</tr>
</tbody>
</table>
| Oral anticoagulants other than VKA  
  - dabigatran  
  - rivaroxaban  
  - apixaban | Lixiana | Discontinue dabigatran, rivaroxaban or apixaban and start Lixiana at the time of the next dose of the oral anticoagulant (see section 5.1). |
| Parenteral anticoagulants | Lixiana | These medicinal products should not be administered simultaneously.  
Subcutaneous anticoagulant (i.e.: LMWH, fondaparinux):  
Discontinue subcutaneous anticoagulant and start Lixiana at the time of the next scheduled subcutaneous anticoagulant dose.  
Intravenous unfractionated heparin (UFH):  
Discontinue the infusion and start Lixiana 4 hours later. |
### Switching from Lixiana

<table>
<thead>
<tr>
<th>From</th>
<th>To</th>
<th>Recommendation</th>
</tr>
</thead>
</table>
| Lixiana                   | Vitamin K Antagonist (VKA)    | There is a potential for inadequate anticoagulation during the transition from Lixiana to VKA. Continuous adequate anticoagulation should be ensured during any transition to an alternate anticoagulant.  

**Oral option:** For patients currently on a 60 mg dose, administer a Lixiana dose of 30 mg once daily together with an appropriate VKA dose.  

For patients currently on a 30 mg dose (for one or more of the following clinical factors: moderate to severe renal impairment (CrCL 15 – 50 mL/min), low body weight, or use with certain P-gp inhibitors), administer a Lixiana dose of 15 mg once daily together with an appropriate VKA dose.  

Patients should not take a loading dose of VKA in order to promptly achieve a stable INR between 2 and 3. It is recommended to take into account the maintenance dose of VKA and if the patient was previously taking a VKA or to use valid INR driven VKA treatment algorithm, in accordance with local practice.  

Once an INR ≥ 2.0 is achieved, Lixiana should be discontinued. Most patients (85%) should be able to achieve an INR ≥ 2.0 within 14 days of concomitant administration of Lixiana and VKA. After 14 days it is recommended that Lixiana is discontinued and the VKA continued to be titrated to achieve an INR between 2 and 3.  

It is recommended that during the first 14 days of concomitant therapy the INR is measured at least 3 times just prior to taking the daily dose of Lixiana to minimise the influence of Lixiana on INR measurements. Concomitant Lixiana and VKA can increase the INR post Lixiana dose by up to 46%.  

**Parenteral option:** Discontinue Lixiana and administer a parenteral anticoagulant and VKA at the time of the next scheduled Lixiana dose. Once a stable INR of ≥ 2.0 is achieved, the parenteral anticoagulant should be discontinued and the VKA continued.  

| Lixiana                   | Oral anticoagulants other than VKA | Discontinue Lixiana and start the non-VKA anticoagulant at the time of the next scheduled dose of Lixiana. |
Switching from Lixiana

<table>
<thead>
<tr>
<th>From</th>
<th>To</th>
<th>Recommendation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lixiana</td>
<td>Parenteral</td>
<td>anticoagulants These agents should not be administered simultaneously.</td>
</tr>
<tr>
<td></td>
<td>anticoagulants</td>
<td>Discontinue Lixiana and start the parenteral anticoagulant at the time of the next</td>
</tr>
<tr>
<td></td>
<td></td>
<td>scheduled dose of Lixiana.</td>
</tr>
</tbody>
</table>

**Special populations**

**Assessment of renal function:**
- Renal function should be assessed in all patients by calculating the creatinine clearance (CrCL) prior to initiation of treatment with Lixiana to exclude patients with end stage renal disease (i.e. CrCL < 15 mL/min), to use the correct Lixiana dose in patients with CrCL 15 – 50 mL/min (30 mg once daily), in patients with CrCL > 50 mL/min (60 mg once daily) and when deciding on the use of Lixiana in patients with increased creatinine clearance (see section 4.4).
- Renal function should also be assessed when a change in renal function is suspected during treatment (e.g. hypovolaemia, dehydration, and in case of concomitant use of certain medicinal products).

The method used to estimate renal function (CrCL in mL/min) during the clinical development of Lixiana was the Cockcroft-Gault method. The formula is as follows:

- For creatinine in µmol/L:
  \[ 1.23 \times (140 - \text{age}) \times \text{weight} \times (0.85 \text{ if female}) \]
  \[ \text{serum creatinine} \times (\mu \text{mol/L}) \]

- For creatinine in mg/dL:
  \[ \frac{(140 - \text{age}) \times \text{weight} \times (0.85 \text{ if female})}{72 \times \text{serum creatinine} \times (\text{mg/dL})} \]

This method is recommended when assessing patients’ CrCL prior to and during Lixiana treatment.

**Renal impairment**
In patients with mild renal impairment (CrCL > 50 – 80 mL/min), the recommended dose is 60 mg Lixiana once daily.

In patients with moderate or severe renal impairment (CrCL 15 – 50 mL/min), the recommended dose is 30 mg Lixiana once daily (see section 5.2).

In patients with end stage renal disease (ESRD) (CrCL < 15 mL/min) or on dialysis, the use of Lixiana is not recommended (see sections 4.4 and 5.2).

**Hepatic impairment**
Lixiana is contraindicated in patients with hepatic disease associated with coagulopathy and clinically relevant bleeding risk (see section 4.3).

In patients with severe hepatic impairment Lixiana is not recommended (see sections 4.4 and 5.2).

In patients with mild to moderate hepatic impairment the recommended dose is 60 mg Lixiana once daily (see section 5.2). Lixiana should be used with caution in patients with mild to moderate hepatic impairment (see section 4.4).
Patients with elevated liver enzymes (ALT/AST > 2 x ULN) or total bilirubin ≥ 1.5 x ULN were excluded in clinical trials. Therefore Lixiana should be used with caution in this population (see sections 4.4 and 5.2). Prior to initiating Lixiana, liver function testing should be performed.

**Body weight**
For patients with body weight ≤ 60 kg, the recommended dose is 30 mg Lixiana once daily (see section 5.2).

**Elderly**
No dose reduction is required (see section 5.2).

**Gender**
No dose reduction is required (see section 5.2).

**Concomitant use of Lixiana with P-glycoprotein (P-gp) inhibitors**
In patients concomitantly taking Lixiana and the following P-gp inhibitors: ciclosporin, dronedarone, erythromycin, or ketoconazole, the recommended dose is 30 mg Lixiana once daily (see section 4.5). No dose reduction is required for concomitant use of amiodarone, quinidine or verapamil (see section 4.5). The use of Lixiana with other P-gp inhibitors including HIV protease inhibitors has not been studied.

**Paediatric population**
The safety and efficacy of Lixiana in children and adolescents less than 18 years of age have not been established. No data are available.

**Patients undergoing cardioversion**
Lixiana can be initiated or continued in patients who may require cardioversion. For transeosophageal echocardiogram (TEE) guided cardioversion in patients not previously treated with anticoagulants, Lixiana treatment should be started at least 2 hours before cardioversion to ensure adequate anticoagulation (see sections 5.1 and 5.2). Cardioversion should be performed no later than 12 hours after the dose of Lixiana on the day of the procedure.

**For all patients undergoing cardioversion:** Confirmation should be sought prior to cardioversion that the patient has taken Lixiana as prescribed. Decisions on initiation and duration of treatment should follow established guidelines for anticoagulant treatment in patients undergoing cardioversion.

**Method of administration**
For oral use.
Lixiana can be taken with or without food (see section 5.2).

### 4.3 Contraindications
- Hypersensitivity to the active substance or to any of the excipients listed in section 6.1.
- Clinically significant active bleeding.
- Hepatic disease associated with coagulopathy and clinically relevant bleeding risk.
- Lesion or condition, if considered to be a significant risk for major bleeding. This may include current or recent gastrointestinal ulceration, presence of malignant neoplasms at high risk of bleeding, recent brain or spinal injury, recent brain, spinal or ophthalmic surgery, recent intracranial haemorrhage, known or suspected oesophageal varices, arteriovenous malformations, vascular aneurysms or major intraspinal or intracerebral vascular abnormalities.
- Uncontrolled severe hypertension.
- Concomitant treatment with any other anticoagulants e.g. unfractionated heparin (UFH), low molecular weight heparins (enoxaparin, dalteparin, etc.), heparin derivatives (fondaparinux, etc.), oral anticoagulants (warfarin, dabigatran etexilate, rivaroxaban, apixaban etc.) except under
specific circumstances of switching oral anticoagulant therapy (see section 4.2) or when UFH is given at doses necessary to maintain an open central venous or arterial catheter (see section 4.5).

- Pregnancy and breast-feeding (see section 4.6).

4.4 Special warnings and precautions for use

Lixiana 15 mg is not indicated as monotherapy, as it may result in decreased efficacy. It is only indicated in the process of switching from Lixiana 30 mg (patients with one or more clinical factors for increased exposure; see table 1) to VKA, together with an appropriate VKA dose (see table 2, section 4.2).

**Haemorrhagic risk**

Edoxaban increases the risk of bleeding and can cause serious, potentially fatal bleeding. Lixiana, like other anticoagulants, is recommended to be used with caution in patients with increased risk of bleeding. Lixiana administration should be discontinued if severe haemorrhage occurs (see sections 4.8 and 4.9).

In the clinical studies mucosal bleedings (e.g. epistaxis, gastrointestinal, genitourinary) and anaemia were seen more frequently during long term edoxaban treatment compared with VKA treatment. Thus, in addition to adequate clinical surveillance, laboratory testing of haemoglobin/haematocrit could be of value to detect occult bleeding, as judged to be appropriate.

Several sub-groups of patients, as detailed below, are at increased risk of bleeding. These patients are to be carefully monitored for signs and symptoms of bleeding complications and anaemia after initiation of treatment (see section 4.8). Any unexplained fall in haemoglobin or blood pressure should lead to a search for a bleeding site.

The anticoagulant effect of edoxaban cannot be reliably monitored with standard laboratory testing. A specific anticoagulant reversal agent for edoxaban is not available (see section 4.9).

Haemodialysis does not significantly contribute to edoxaban clearance (see section 5.2).

**Elderly**

The co-administration of Lixiana with ASA in elderly patients should be used cautiously because of a potentially higher bleeding risk (see section 4.5).

**Renal impairment**

The plasma AUC for subjects with mild (CrCL > 50 - 80 mL/min), moderate (CrCL 30 - 50 mL/min) and severe (CrCL < 30 mL/min but not undergoing dialysis) renal impairment was increased by 32%, 74%, and 72%, respectively, relative to subjects with normal renal function (see section 4.2 for dose reduction).

In patients with end stage renal disease or on dialysis, Lixiana is not recommended (see sections 4.2 and 5.2).

**Renal function in NVAF**

A trend towards decreasing efficacy with increasing creatinine clearance was observed for edoxaban compared to well-managed warfarin (see section 5.1). Therefore, edoxaban should only be used in patients with NVAF and high creatinine clearance after a careful evaluation of the individual thromboembolic and bleeding risk.

Assessment of renal function: CrCL should be monitored at the beginning of the treatment in all patients and afterwards when clinically indicated (see section 4.2).

**Hepatic impairment**

Lixiana is not recommended in patients with severe hepatic impairment (see sections 4.2 and 5.2).
Lixiana should be used with caution in patients with mild or moderate hepatic impairment (see section 4.2).

Patients with elevated liver enzymes (ALT/AST > 2 x ULN) or total bilirubin ≥ 1.5 x ULN were excluded in clinical trials. Therefore Lixiana should be used with caution in this population (see sections 4.2 and 5.2). Prior to initiating Lixiana, liver function testing should be performed. Periodic hepatic monitoring is recommended for patients on Lixiana treatment beyond 1 year.

**Discontinuation for surgery and other interventions**

If anticoagulation must be discontinued to reduce the risk of bleeding with surgical or other procedures, Lixiana should be stopped as soon as possible and preferably at least 24 hours before the procedure.

In deciding whether a procedure should be delayed until 24 hours after the last dose of Lixiana, the increased risk of bleeding should be weighed against the urgency of the intervention. Lixiana should be restarted after the surgical or other procedures as soon as adequate haemostasis has been established, noting that the time to onset of the edoxaban anticoagulant therapeutic effect is 1 – 2 hours. If oral medicinal products cannot be taken during or after surgical intervention, consider administering a parenteral anticoagulant and then switch to oral once daily Lixiana (see section 4.2).

**Anticoagulants, antiplatelets, and thrombolytics**

Concomitant use of medicines affecting haemostasis may increase the risk of bleeding. These include acetylsalicylic acid (ASA), P2Y12 platelet inhibitors, other antithrombotic agents, fibrinolytic therapy, and chronic nonsteroidal anti-inflammatory drugs (NSAIDs) (see section 4.5).

**Prosthetic heart valves and moderate to severe mitral stenosis**

Edoxaban has not been studied in patients with mechanical heart valves, in patients during the first 3 months after implantation of a bioprosthetic heart valve, with or without atrial fibrillation, or in patients with moderate to severe mitral stenosis. Therefore, use of edoxaban is not recommended in these patients.

**Haemodynamically unstable PE patients or patients who require thrombolysis or pulmonary embolectomy**

Lixiana is not recommended as an alternative to unfractionated heparin in patients with pulmonary embolism who are haemodynamically unstable or may receive thrombolysis or pulmonary embolectomy since the safety and efficacy of edoxaban have not been established in these clinical situations.

**Patients with active cancer**

Efficacy and safety of edoxaban in the treatment and/or prevention of VTE in patients with active cancer have not been established.

**Laboratory coagulation parameters**

Although treatment with edoxaban does not require routine monitoring, the effect on anticoagulation can be estimated by a calibrated quantitative anti-Factor Xa assay which may help to inform clinical decisions in particular situations as, e.g. overdose and emergency surgery (see also section 5.2).

Edoxaban prolongs standard clotting tests such as prothrombin time (PT), INR, and activated partial thromboplastin time (aPTT) as a result of FXa inhibition. Changes observed in these clotting tests at the expected therapeutic dose are, however, small, subject to a high degree of variability, and not useful in monitoring the anticoagulation effect of edoxaban.
4.5 Interaction with other medicinal products and other forms of interaction

Edoxaban is predominantly absorbed in the upper gastrointestinal (GI) tract. Thus, medicines or disease conditions that increase gastric emptying and gut motility have the possibility of reducing edoxaban dissolution and absorption.

**P-gp inhibitors**

Edoxaban is a substrate for the efflux transporter P-gp. In pharmacokinetic (PK) studies, concomitant administration of edoxaban with the P-gp inhibitors: ciclosporin, dronedarone, erythromycin, ketoconazole, quinidine, or verapamil resulted in increased plasma concentrations of edoxaban. Concomitant use of edoxaban with ciclosporin, dronedarone, erythromycin, or ketoconazole requires dose reduction to 30 mg once daily. Concomitant use of edoxaban with quinidine, verapamil, or amiodarone does not require dose reduction based on clinical data (see section 4.2).

The use of edoxaban with other P-gp inhibitors including HIV protease inhibitors has not been studied.

Lixiana 30 mg once daily must be administered during concomitant use with the following P-gp inhibitors:

- **Ciclosporin**: Concurrent administration of a single dose of ciclosporin 500 mg with a single dose of edoxaban 60 mg increased edoxaban AUC and C\text{max} by 73% and 74%, respectively.
- **Dronedarone**: Dronedarone 400 mg twice daily for 7 days with a single concomitant dose of edoxaban 60 mg on Day 5 increased edoxaban AUC and C\text{max} by 85% and 46%, respectively.
- **Erythromycin**: Erythromycin 500 mg four times daily for 8 days with a single concomitant dose of edoxaban 60 mg on Day 7 increased the edoxaban AUC and C\text{max} by 85% and 68%, respectively.
- **Ketoconazole**: Ketoconazole 400 mg once daily for 7 days with a single concomitant dose of edoxaban 60 mg on Day 4, increased edoxaban AUC and C\text{max} by 87% and 89%, respectively.

Lixiana 60 mg once daily is recommended during concomitant use with the following P-gp inhibitors:

- **Quinidine**: Quinidine 300 mg once daily on Days 1 and 4 and three times daily on Days 2 and 3, with a single concomitant dose of edoxaban 60 mg on Day 3, increased edoxaban AUC over 24 hours by 77% and C\text{max} by 85%, respectively.
- **Verapamil**: Verapamil 240 mg once daily for 11 days with a single concomitant dose of edoxaban 60 mg on Day 10 increased the edoxaban AUC and C\text{max} by approximately 53%.
- **Amiodarone**: Co-administration of amiodarone 400 mg once daily with edoxaban 60 mg once daily increased AUC by 40% and C\text{max} by 66%. This was not considered clinically significant. In ENGAGE AF-TIMI 48 study in NVAF, efficacy and safety results were similar for subjects with and without concomitant amiodarone use.

**P-gp inducers**

Co-administration of edoxaban with the P-gp inducer rifampicin led to a decrease in mean edoxaban AUC and a shortened half-life, with possible decreases in its pharmacodynamic effects. The concomitant use of edoxaban with other P-gp inducers (e.g. phenytoin, carbamazepine, phenobarbital or St. John's Wort) may lead to reduced edoxaban plasma concentrations. Edoxaban should be used with caution when co-administered with P-gp inducers.

**P-gp substrates**

**Digoxin**: Edoxaban 60 mg once daily on days 1 to 14 with coadministration of multiple daily doses of digoxin 0.25 mg twice daily (days 8 and 9) and 0.25 mg once daily (days 10 to 14) increased the C\text{max} of edoxaban by 17%, with no significant effect on AUC or renal clearance at steady state. When the effects of edoxaban on digoxin PK were also examined, the C\text{max} of digoxin increased by approximately 28% and AUC by 7%. This was not considered clinically relevant. No dose modification is necessary when Lixiana is administered with digoxin.
**Anticoagulants, antiplatelets, and NSAIDs**

**Anticoagulants:** Co-administration of edoxaban with other anticoagulants is contraindicated due to increased risk of bleeding (see section 4.3).

**Acetylsalicylic acid (ASA):** Co-administration of ASA (100 mg or 325 mg) and edoxaban increased bleeding time relative to either medicine alone. Co-administration of high dose ASA (325 mg) increased the steady state C\text{max} and AUC of edoxaban by 35% and 32%, respectively. The concomitant chronic use of high dose ASA (325 mg) with edoxaban is not recommended. Concomitant administration of higher doses than 100 mg ASA should only be performed under medical supervision.

In clinical studies concomitant use of ASA (low dose ≤ 100 mg/day), other antiplatelet agents, and thienopyridines was permitted and resulted in approximately a 2-fold increase in major bleeding in comparison with no concomitant use, although to a similar extent in the edoxaban and warfarin groups (see section 4.4). Co-administration of low dose ASA (≤ 100 mg) did not affect the peak or total exposure of edoxaban either after single dose or at steady-state. Edoxaban can be co-administered with low dose ASA (≤ 100 mg/day).

**Platelet inhibitors:** In ENGAGE AF-TIMI 48 concomitant use of thienopyridines (e.g. clopidogrel) monotherapy was permitted and resulted in increased clinically relevant bleeding although with a lower risk of bleeding on edoxaban compared to warfarin (see section 4.4).

There is very limited experience on the use of edoxaban with dual antiplatelet therapy or fibrinolytic agents.

**NSAIDs:** Co-administration of naproxen and edoxaban increased bleeding time relative to either medicine alone. Naproxen had no effect on the C\text{max} and AUC of edoxaban. In clinical studies, co-administration of NSAIDs resulted in increased clinically relevant bleeding. Chronic use of NSAIDs with edoxaban is not recommended.

**Effect of edoxaban on other medicines**

Edoxaban increased the C\text{max} of concomitantly administered digoxin by 28%; however, the AUC was not affected. Edoxaban had no effect on the C\text{max} and AUC of quinidine.

Edoxaban decreased the C\text{max} and AUC of concomitantly administered verapamil by 14% and 16%, respectively.

### 4.6 Fertility, pregnancy and lactation

**Woman of childbearing potential**

Women of childbearing potential should avoid becoming pregnant during treatment with edoxaban.

**Pregnancy**

Safety and efficacy of edoxaban have not been established in pregnant women. Studies in animals have shown reproductive toxicity (see section 5.3). Due to the potential reproductive toxicity, the intrinsic risk of bleeding and the evidence that edoxaban passes the placenta, Lixiana is contraindicated during pregnancy (see section 4.3).

**Breast-feeding**

Safety and efficacy of edoxaban have not been established in breast-feeding women. Data from animals indicate that edoxaban is secreted into breast milk. Therefore Lixiana is contraindicated during breast-feeding (see section 4.3). A decision must be made whether to discontinue breast-feeding or to discontinue/abstain from therapy.

**Fertility**

No specific studies with edoxaban in humans have been conducted to evaluate effects on fertility. In a study on male and female fertility in rats no effects were seen (see section 5.3).
4.7 Effects on ability to drive and use machines

Lixiana has no or negligible influence on the ability to drive and use machines.

4.8 Undesirable effects

Summary of the safety profile

The safety of edoxaban has been evaluated in two Phase 3 studies including 21,105 patients with NVAF (ENGAGE AF-TIMI 48 study), and 8,292 patients with VTE (DVT and PE) (Hokusai-VTE study).

The average exposure to edoxaban 60 mg (including 30 mg dose reduced) was 2.5 years among 7,012 patients in ENGAGE AF-TIMI 48 and 251 days among 4,118 patients in Hokusai-VTE. Adverse reactions were experienced by 2,256 (32.2%) of the patients treated with edoxaban 60 mg (30 mg dose reduced) in the ENGAGE AF-TIMI 48 study and 1,249 (30.3%) in the Hokusai-VTE study.

In both studies, the most common adverse reactions related to bleeding with edoxaban 60 mg based on adjudicated terms included cutaneous soft tissue haemorrhage (up to 5.9%) and epistaxis (up to 4.7%), while vaginal haemorrhage (9.0%) was the most common bleeding-related adverse reaction in Hokusai-VTE only.

Bleeding can occur at any site and may be severe and even fatal (see section 4.4). Common other adverse reactions for edoxaban were anaemia, rash and abnormal liver function tests.

Tabulated list of adverse reactions

Table 3 provides the list of adverse reactions from the two pivotal Phase 3 studies in patients with VTE (DVT and PE) (Hokusai-VTE study) and AF (ENGAGE AF-TIMI 48 study) combined for both indications. The adverse reactions are classified by System Organ Class and frequency, using the following convention:

Very common (≥ 1/10), Common (≥ 1/100 to < 1/10), Uncommon (≥ 1/1,000 to < 1/100), Rare (≥ 1/10,000 to < 1/100), Very rare (< 1/10,000), not known (cannot be estimated from the available data).

**Table 3: List of adverse reactions for NVAF and VTE**

<table>
<thead>
<tr>
<th>System Organ Class</th>
<th>Frequency</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Blood and lymphatic system disorders</strong></td>
<td></td>
</tr>
<tr>
<td>Anaemia</td>
<td>Common</td>
</tr>
<tr>
<td>Thrombocytopenia</td>
<td>Uncommon</td>
</tr>
<tr>
<td><strong>Immune system disorders</strong></td>
<td></td>
</tr>
<tr>
<td>Hypersensitivity</td>
<td>Uncommon</td>
</tr>
<tr>
<td>Anaphylactic reaction</td>
<td>Rare</td>
</tr>
<tr>
<td>Allergic oedema</td>
<td>Rare</td>
</tr>
<tr>
<td><strong>Nervous system disorders</strong></td>
<td></td>
</tr>
<tr>
<td>Dizziness</td>
<td>Common</td>
</tr>
<tr>
<td>Headache</td>
<td>Common</td>
</tr>
<tr>
<td>Intracranial haemorrhage (ICH)</td>
<td>Uncommon</td>
</tr>
<tr>
<td>Subarachnoid haemorrhage</td>
<td>Rare</td>
</tr>
<tr>
<td><strong>Eye disorders</strong></td>
<td></td>
</tr>
<tr>
<td>Conjunctival/Scleral haemorrhage</td>
<td>Uncommon</td>
</tr>
<tr>
<td>Intraocular haemorrhage</td>
<td>Uncommon</td>
</tr>
<tr>
<td><strong>Cardiac disorders</strong></td>
<td></td>
</tr>
<tr>
<td>Pericardial haemorrhage</td>
<td>Rare</td>
</tr>
<tr>
<td>System Organ Class</td>
<td>Frequency</td>
</tr>
<tr>
<td>-------------------</td>
<td>-----------</td>
</tr>
<tr>
<td><strong>Vascular disorders</strong></td>
<td></td>
</tr>
<tr>
<td>Other haemorrhage</td>
<td>Uncommon</td>
</tr>
<tr>
<td><strong>Respiratory, thoracic and mediastinal disorders</strong></td>
<td></td>
</tr>
<tr>
<td>Epistaxis</td>
<td>Common</td>
</tr>
<tr>
<td>Haemoptysis</td>
<td>Uncommon</td>
</tr>
<tr>
<td><strong>Gastrointestinal disorders</strong></td>
<td></td>
</tr>
<tr>
<td>Abdominal pain</td>
<td>Common</td>
</tr>
<tr>
<td>Lower GI haemorrhage</td>
<td>Common</td>
</tr>
<tr>
<td>Upper GI haemorrhage</td>
<td>Common</td>
</tr>
<tr>
<td>Oral/Pharyngeal haemorrhage</td>
<td>Common</td>
</tr>
<tr>
<td>Nausea</td>
<td>Common</td>
</tr>
<tr>
<td>Retroperitoneal haemorrhage</td>
<td>Rare</td>
</tr>
<tr>
<td><strong>Hepatobiliary disorders</strong></td>
<td></td>
</tr>
<tr>
<td>Blood bilirubin increased</td>
<td>Common</td>
</tr>
<tr>
<td>Gammaglutamyltransferase increased</td>
<td>Common</td>
</tr>
<tr>
<td>Blood alkaline phosphatase increased</td>
<td>Uncommon</td>
</tr>
<tr>
<td>Transaminases increased</td>
<td>Uncommon</td>
</tr>
<tr>
<td>Aspartate aminotransferase increased</td>
<td>Uncommon</td>
</tr>
<tr>
<td><strong>Skin and subcutaneous tissue disorders</strong></td>
<td></td>
</tr>
<tr>
<td>Cutaneous soft tissue haemorrhage</td>
<td>Common</td>
</tr>
<tr>
<td>Rash</td>
<td>Common</td>
</tr>
<tr>
<td>Pruritus</td>
<td>Common</td>
</tr>
<tr>
<td>Urticaria</td>
<td>Common</td>
</tr>
<tr>
<td><strong>Musculoskeletal and connective tissue disorders</strong></td>
<td></td>
</tr>
<tr>
<td>Intramuscular haemorrhage (no compartment syndrome)</td>
<td>Rare</td>
</tr>
<tr>
<td>Intra-articular haemorrhage</td>
<td>Rare</td>
</tr>
<tr>
<td><strong>Renal and urinary disorders</strong></td>
<td></td>
</tr>
<tr>
<td>Macroscopic haematuria/urethral haemorrhage</td>
<td>Common</td>
</tr>
<tr>
<td><strong>Reproductive system and breast disorders</strong></td>
<td></td>
</tr>
<tr>
<td>Vaginal haemorrhage(^1)</td>
<td>Common</td>
</tr>
<tr>
<td><strong>General disorders and administration site conditions</strong></td>
<td></td>
</tr>
<tr>
<td>Puncture site haemorrhage</td>
<td>Common</td>
</tr>
<tr>
<td><strong>Investigations</strong></td>
<td></td>
</tr>
<tr>
<td>Liver function test abnormal</td>
<td>Common</td>
</tr>
<tr>
<td><strong>Injury, poisoning and procedural complications</strong></td>
<td></td>
</tr>
<tr>
<td>Surgical site haemorrhage</td>
<td>Uncommon</td>
</tr>
<tr>
<td>Subdural haemorrhage</td>
<td>Rare</td>
</tr>
<tr>
<td>Procedural haemorrhage</td>
<td>Rare</td>
</tr>
</tbody>
</table>

\(^1\) Reporting rates are based on the female population in clinical trials. Vaginal bleeds were reported commonly in women under the age of 50 years, while it was uncommon in women over the age of 50 years.
Description of selected adverse reactions

Due to the pharmacological mode of action, the use of Lixiana may be associated with an increased risk of occult or overt bleeding from any tissue or organ which may result in post haemorrhagic anaemia. The signs, symptoms, and severity (including fatal outcome) will vary according to the location and degree or extent of the bleeding and/or anaemia (see section 4.9 Management of bleeding). In the clinical studies mucosal bleedings (e.g. epistaxis, gastrointestinal, genitourinary) and anaemia were seen more frequently during long term edoxaban treatment compared with VKA treatment. Thus, in addition to adequate clinical surveillance, laboratory testing of haemoglobin/haematocrit could be of value to detect occult bleeding, as judged to be appropriate. The risk of bleedings may be increased in certain patient groups e.g. those patients with uncontrolled severe arterial hypertension and/or on concomitant treatment affecting haemostasis (see Haemorrhagic risk in section 4.4). Menstrual bleeding may be intensified and/or prolonged. Haemorrhagic complications may present as weakness, paleness, dizziness, headache or unexplained swelling, dyspnoea, and unexplained shock.

Known complications secondary to severe bleeding such as compartment syndrome and renal failure due to hypoperfusion have been reported for Lixiana. Therefore, the possibility of haemorrhage is to be considered in evaluating the condition in any anticoagulated patient.

Reporting of suspected adverse reactions

Reporting suspected adverse reactions after authorisation of the medicinal product is important. It allows continued monitoring of the benefit/risk balance of the medicinal product. Healthcare professionals are asked to report any suspected adverse reactions via the national reporting system listed in Appendix V.

4.9 Overdose

Overdose with edoxaban may lead to haemorrhage. Experience with overdose cases is very limited.

A specific antidote antagonising the pharmacodynamic effect of edoxaban is not available.

Early administration of activated charcoal may be considered in case of edoxaban overdose to reduce absorption. This recommendation is based on standard treatment of drug overdose and data available with similar compounds, as the use of activated charcoal to reduce absorption of edoxaban has not been specifically studied in the edoxaban clinical programme.

Management of bleeding

Should a bleeding complication arise in a patient receiving edoxaban, the next edoxaban administration should be delayed or treatment should be discontinued as appropriate. Edoxaban has a half-life of approximately 10 to 14 hours (see section 5.2). Management should be individualised according to the severity and location of the haemorrhage. Appropriate symptomatic treatment could be used as needed, such as mechanical compression (e.g. for severe epistaxis), surgical haemostasis with bleeding control procedures, fluid replacement and haemodynamic support, blood products (packed red cells or fresh frozen plasma, depending on associated anaemia or coagulopathy) or platelets.

For life-threatening bleeding that cannot be controlled with the measures such as transfusion or haemostasis, the administration of a 4-factor prothrombin complex concentrate (PCC) at 50 IU/kg has been shown to reverse the effects of Lixiana 30 minutes after completing the infusion.

Recombinant factor VIIa (r-FVIIa) can also be considered. However, there is limited clinical experience with the use of this product in individuals receiving edoxaban.

Depending on local availability, a consultation with a coagulation expert should be considered in case of major bleedings.

Protamine sulfate and vitamin K are not expected to affect the anticoagulant activity of edoxaban.
There is no experience with antifibrinolytic agents (tranexamic acid, aminocaproic acid) in individuals receiving edoxaban. There is neither scientific rationale for benefit nor experience with the use of systemic haemostatics (desmopressin, aprotinin) in individuals receiving edoxaban. Due to the high plasma protein binding edoxaban is not expected to be dialysable.

5. PHARMACOLOGICAL PROPERTIES

5.1 Pharmacodynamic properties

Pharmacotherapeutic group: Other antithrombotic agents, ATC code: B01AF03

Mechanism of action
Edoxaban is a highly selective, direct and reversible inhibitor of factor Xa, the serine protease located in the final common pathway of the coagulation cascade. Edoxaban inhibits free factor Xa, and prothrombinase activity. Inhibition of factor Xa in the coagulation cascade reduces thrombin generation, prolongs clotting time and reduces the risk of thrombus formation.

Pharmacodynamic effects
Edoxaban produces rapid onset of pharmacodynamic effects within 1 - 2 hours, which corresponds with peak edoxaban exposure (Cmax). The pharmacodynamic effects measured by anti-factor Xa assay are predictable and correlate with the dose and the concentration of edoxaban. As a result of FXa inhibition, edoxaban also prolongs clotting time in tests such as prothrombin time (PT), and activated partial thromboplastin time (aPTT). Changes observed in these clotting tests are expected at the therapeutic dose, however, these changes are small, subject to a high degree of variability, and not useful in monitoring the anticoagulation effect of edoxaban.

Effects of coagulation markers when switching from rivaroxaban, dabigatran, or apixaban to edoxaban
In clinical pharmacology studies, healthy subjects received rivaroxaban 20 mg once daily, dabigatran 150 mg twice daily, or apixaban 5 mg twice daily, followed by a single dose of edoxaban 60 mg on Day 4. The effect on prothrombin time (PT) and other coagulation biomarkers (e.g. anti-FXa, aPTT) was measured. Following the switch to edoxaban on Day 4 the PT was equivalent to Day 3 of rivaroxaban and apixaban. For dabigatran higher aPTT activity was observed after edoxaban administration with prior dabigatran treatment compared to that after treatment with edoxaban alone. This is considered to be due to the carry-over effect of dabigatran treatment, however, this did not lead to a prolongation of bleeding time.
Based on these data, when switching from these anticoagulants to edoxaban, the first dose of edoxaban can be initiated at the time of the next scheduled dose of the previous anticoagulant (see section 4.2).

Clinical efficacy and safety

Prevention of stroke and systemic embolism
The edoxaban clinical programme for atrial fibrillation was designed to demonstrate the efficacy and safety of two dose groups of edoxaban compared to warfarin for the prevention of stroke and systemic embolism in subjects with nonvalvular atrial fibrillation and at moderate to high risk of stroke and systemic embolic events (SEE).

In the pivotal ENGAGE AF-TIMI 48 study (an event-driven, Phase 3, multi-centre, randomised, double-blind double-dummy parallel-group study), 21,105 subjects, with a mean CHADS2 score of 2.8, were randomised to either edoxaban 30 mg once daily treatment group, or edoxaban 60 mg once daily treatment group or warfarin. Subjects in both edoxaban treatment groups had their dose halved if one or more of the following clinical factors were present: moderate renal impairment (CrCL 30 – 50 mL/min), low body weight (≤ 60 kg) or concomitant use of specific P-gp inhibitors (verapamil, quinidine, dronedarone).
The primary efficacy endpoint was the composite of stroke and SEE. Secondary efficacy endpoints included: Composite of stroke, SEE, and cardiovascular (CV) mortality; major adverse cardiovascular event (MACE), which is the composite of non-fatal MI, non-fatal stroke, non-fatal SEE, and death due to CV cause or bleeding; composite of stroke, SEE, and all-cause mortality.

The median study drug exposure for both the edoxaban 60 mg and 30 mg treatment groups was 2.5 years. The median study follow-up for both the edoxaban 60 mg and 30 mg treatment groups was 2.8 years. The median subject-year exposure was 15,471, and 15,840 for the 60 mg and 30 mg treatment groups, respectively; and the median subject-year follow-up was 19,191 and 19,216 for the 60 mg and 30 mg treatment groups, respectively.

In the warfarin group, the median TTR (time in therapeutic range, INR 2.0 to 3.0) was 68.4%.

The main analysis of efficacy was aimed to show the non-inferiority of edoxaban versus warfarin on first stroke or SEE that occurred during treatment or within 3 days from the last dose taken in the modified intention-to-treat (mITT) population. Edoxaban 60 mg was non-inferior to warfarin for the primary efficacy endpoint of stroke or SEE (upper limit of the 97.5% CI of the HR was below the pre-specified non-inferiority margin of 1.38) (Table 4).

Table 4: Strokes and Systemic Embolic Events in the ENGAGE AF–TIMI 48 Study - mITT, on-treatment

<table>
<thead>
<tr>
<th>Primary Endpoint</th>
<th>Edoxaban 60 mg (30 mg Dose Reduced) (N = 7,012)</th>
<th>Warfarin (N = 7,012)</th>
</tr>
</thead>
<tbody>
<tr>
<td>First Stroke/SEE</td>
<td></td>
<td></td>
</tr>
<tr>
<td>n</td>
<td>182</td>
<td>232</td>
</tr>
<tr>
<td>Event Rate (%/yr)</td>
<td>1.18</td>
<td>1.50</td>
</tr>
<tr>
<td>HR (97.5% CI)</td>
<td>0.79 (0.63, 0.99)</td>
<td></td>
</tr>
<tr>
<td>p-value for non-inferiority&lt;sup&gt;c&lt;/sup&gt;</td>
<td>&lt;0.0001</td>
<td></td>
</tr>
<tr>
<td>First Ischaemic Stroke</td>
<td></td>
<td></td>
</tr>
<tr>
<td>n</td>
<td>135</td>
<td>144</td>
</tr>
<tr>
<td>Event Rate (%/yr)</td>
<td>0.87</td>
<td>0.93</td>
</tr>
<tr>
<td>HR (95% CI)</td>
<td>0.94 (0.75, 1.19)</td>
<td></td>
</tr>
<tr>
<td>First Haemorrhagic Stroke</td>
<td></td>
<td></td>
</tr>
<tr>
<td>n</td>
<td>40</td>
<td>76</td>
</tr>
<tr>
<td>Event Rate (%/yr)</td>
<td>0.26</td>
<td>0.49</td>
</tr>
<tr>
<td>HR (95% CI)</td>
<td>0.53 (0.36, 0.78)</td>
<td></td>
</tr>
<tr>
<td>First SEE</td>
<td></td>
<td></td>
</tr>
<tr>
<td>n (%/yr)&lt;sup&gt;a&lt;/sup&gt;</td>
<td>8 (0.05)</td>
<td>13 (0.08)</td>
</tr>
<tr>
<td>HR (95% CI)</td>
<td>0.62 (0.26, 1.50)</td>
<td></td>
</tr>
</tbody>
</table>

Abbreviations: HR = Hazard Ratio versus warfarin, CI = Confidence Interval, n = number of events, mITT = modified Intent To Treat, N = number of subjects in mITT population, SEE = Systemic Embolic Event, yr = year.

<sup>a</sup> A subject can be represented in multiple rows.

<sup>b</sup> The event rate (%/yr) is calculated as number of events/subject-year exposure.

<sup>c</sup> The two-sided p-value is based on the non-inferiority margin of 1.38.

During the overall study period in the ITT population (analysis set to show superiority), adjudicated stroke or SEE occurred in 296 subjects in the edoxaban 60 mg group (1.57% per year), and 337
subjects in the warfarin group (1.80% per year). Compared to warfarin-treated subjects, the HR in the edoxaban 60 mg group was 0.87 (99% CI: 0.71, 1.07, p = 0.08 for superiority).

In subgroup analyses, for subjects in the 60 mg treatment group who were dose reduced to 30 mg in the ENGAGE AF-TIMI 48 study (for body weight ≤ 60 kg, moderate renal impairment, or concomitant use of P-gp inhibitors), the event rate was: 2.29% per year for the primary endpoint, compared to the event rate of 2.66% per year for the matching subjects in the warfarin group [HR (95% CI): 0.86 (0.66, 1.13)].

The efficacy results for pre-specified major subgroups (with dose reduction as required), including age, body weight, gender, status of renal function, prior stroke or TIA, diabetes and P-gp inhibitors were generally consistent with the primary efficacy results for the overall population studied in the trial.

The Hazard Ratio (Edoxaban 60 mg vs. warfarin) for the primary endpoint in the centres with a lower average time of INR in the target range (INR TTR) for warfarin was 0.73 – 0.80 for the lowest 3 quartiles (INR TTR ≤ 57.7% to ≤ 73.9%). It was 1.07 in centres with the best control of warfarin therapy (4th quartile with > 73.9% of INR values in the therapeutic range).

There was a statistically significant interaction between the effect of edoxaban versus warfarin on the main study outcome (stroke/SEE) and renal function (p-value 0.0042; mITT, overall study period).

Table 5 shows ischaemic strokes/SEE by creatinine clearance category in NVAF patients in ENGAGE AF-TIMI 48. There is a decreasing event rate at increasing CrCL in both treatment groups.

Table 5: Number of Ischaemic Strokes/SEE by creatinine clearance category in ENGAGE AF-TIMI 48, mITT Analysis Set Overall Study

<table>
<thead>
<tr>
<th>CrCL subgroup (mL/min)</th>
<th>Edoxaban 60 mg (N = 7,012)</th>
<th>Warfarin (N = 7,012)</th>
</tr>
</thead>
<tbody>
<tr>
<td>n</td>
<td>Number of Events</td>
<td>Event rate (%/year)</td>
</tr>
<tr>
<td>≥ 30 to ≤ 50</td>
<td>1,302</td>
<td>63</td>
</tr>
<tr>
<td>&gt; 50 to ≤ 70</td>
<td>2,093</td>
<td>85</td>
</tr>
<tr>
<td>&gt; 70 to ≤ 90</td>
<td>1,661</td>
<td>45</td>
</tr>
<tr>
<td>&gt; 90 to ≤ 110</td>
<td>927</td>
<td>27</td>
</tr>
<tr>
<td>&gt; 110 to ≤ 130</td>
<td>497</td>
<td>14</td>
</tr>
<tr>
<td>&gt; 130</td>
<td>462</td>
<td>10</td>
</tr>
</tbody>
</table>

Abbreviations: N = number of subjects; mITT population overall study period; n = number of patients in subgroup
*HR not computed if number of events < 5 in one treatment group.

Within renal function subgroups, results for the secondary efficacy endpoints were consistent with those for the primary endpoint.

Superiority testing was performed on the ITT Overall Study Period. Stroke and SEE occurred in fewer subjects in the edoxaban 60 mg treatment group than in the warfarin group (1.57% and 1.80% per year, respectively), with a HR of 0.87 (99% CI: 0.71, 1.07, p = 0.0807 for superiority).
The pre-specified composite endpoints for the comparison of the edoxaban 60 mg treatment group to warfarin for stroke, SEE, and CV mortality HR (99% CI) was 0.87 (0.76, 0.99), MACE 0.89 (0.78, 1.00), and stroke, SEE, and all-cause mortality 0.90 (0.80, 1.01).

The results for all-cause mortality ( adjudicated deaths) in the ENGAGE AF-TIMI 48 study were 769 (3.99% per year) for subjects taking edoxaban 60 mg (30 mg dose reduced) as opposed to 836 (4.35% per year) for warfarin [HR (95% CI): 0.91 (0.83, 1.01)].

All-cause mortality (adjudicated deaths) per renal subgroups (edoxaban vs. warfarin):
- CrCL 30 to ≤ 50 mL/min [HR (95% CI): 0.81 (0.68, 0.97)];
- CrCL > 50 to < 80 mL/min [HR (95% CI): 0.87 (0.75, 1.02)];
- CrCL ≥ 80 mL/min [HR (95% CI): 1.15 (0.95, 1.40)].

Edoxaban 60 mg (30 mg dose reduced) resulted in a lower rate of cardiovascular mortality compared to warfarin [HR (95% CI): 0.86 (0.77, 0.97)].

Adjudicated efficacy cardiovascular mortality per renal subgroups (edoxaban vs. warfarin):
- CrCL 30 to ≤ 50 mL/min [HR (95% CI): 0.80 (0.65, 0.99)];
- CrCL > 50 to < 80 mL/min [HR (95% CI): 0.75 (0.62, 0.90)];
- CrCL ≥ 80 mL/min [HR (95% CI): 1.16 (0.92, 1.46)].

Safety in patients with NVAF in ENGAGE AF-TIMI 48

The primary safety endpoint was major bleeding.

There was a significant risk reduction in favour of the edoxaban 60 mg treatment group compared with the warfarin group in major bleeding (2.75%, and 3.43% per year, respectively) [HR (95% CI): 0.80 (0.71, 0.91); p = 0.0009], ICH (0.39%, and 0.85% per year, respectively) [HR (95% CI): 0.47 (0.34, 0.63); p < 0.0001], and other types of bleeding (Table 6).

The reduction in fatal bleeds was also significant for the edoxaban 60 mg treatment group compared with the warfarin group (0.21%, and 0.38%) [HR (95% CI): 0.55 (0.36, 0.84); p = 0.0059 for superiority], primarily because of the reduction in fatal ICH bleeds [HR (95% CI): 0.58 (0.35, 0.95); p = 0.0312].
Table 6: Bleeding Events in ENGAGE AF-TIMI 48 Study - Safety Analysis On-Treatment

<table>
<thead>
<tr>
<th></th>
<th>Edoxaban 60 mg (30 mg Dose Reduced) (N = 7,012)</th>
<th>Warfarin (N = 7,012)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Major Bleeding</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>n</td>
<td>418</td>
<td>524</td>
</tr>
<tr>
<td>Event rate (%/yr)a</td>
<td>2.75</td>
<td>3.43</td>
</tr>
<tr>
<td>HR (95% CI)</td>
<td>0.80 (0.71, 0.91)</td>
<td></td>
</tr>
<tr>
<td>p-value</td>
<td>0.0009</td>
<td></td>
</tr>
<tr>
<td>ICHb</td>
<td></td>
<td></td>
</tr>
<tr>
<td>n</td>
<td>61</td>
<td>132</td>
</tr>
<tr>
<td>Event rate (%/yr)a</td>
<td>0.39</td>
<td>0.85</td>
</tr>
<tr>
<td>HR (95% CI)</td>
<td>0.47 (0.34, 0.63)</td>
<td></td>
</tr>
<tr>
<td><strong>Fatal Bleeding</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>n</td>
<td>32</td>
<td>59</td>
</tr>
<tr>
<td>Event rate (%/yr)a</td>
<td>0.21</td>
<td>0.38</td>
</tr>
<tr>
<td>HR (95% CI)</td>
<td>0.55 (0.36, 0.84)</td>
<td></td>
</tr>
<tr>
<td><strong>CRNM Bleeding</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>n</td>
<td>1,214</td>
<td>1,396</td>
</tr>
<tr>
<td>Event rate (%/yr)a</td>
<td>8.67</td>
<td>10.15</td>
</tr>
<tr>
<td>HR (95% CI)</td>
<td>0.86 (0.80, 0.93)</td>
<td></td>
</tr>
<tr>
<td><strong>Any Confirmed Bleedingc</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>n</td>
<td>1,865</td>
<td>2,114</td>
</tr>
<tr>
<td>Event rate (%/yr)a</td>
<td>14.15</td>
<td>16.40</td>
</tr>
<tr>
<td>HR (95% CI)</td>
<td>0.87 (0.82, 0.92)</td>
<td></td>
</tr>
</tbody>
</table>

Abbreviations: ICH = Intracranial Haemorrhage, HR = Hazard Ratio versus warfarin, CI = Confidence Interval, CRNM = Clinically Relevant Non-Major, n = number of subjects with events, N = number of subjects in Safety population, yr = year.

- The event rate (%/yr) is calculated as number of events/subject-year exposure.
- ICH includes primary haemorrhagic stroke, subarachnoid haemorrhage, epi-/subdural haemorrhage, and ischaemic stroke with major haemorrhagic conversion. All ICHs reported on the Adjudicated Cerebrovascular and Non-Intracranial bleed eCRF forms confirmed by the adjudicators are included in ICH counts.
- 'Any Confirmed Bleeding' includes those that the adjudicator defined as clinically overt.

Note: A subject can be included in multiple sub-categories if he/she had an event for those categories. The first event of each category is included in the analysis.

Tables 7, 8 and 9 show major, fatal and intracranial bleedings, respectively, by creatinine clearance category in NVAF patients in ENGAGE AF-TIMI 48. There is a decreasing event rate at increasing CrCL in both treatment groups.
Table 7: Number of Major Bleeding Events by creatinine clearance category in ENGAGE AF-TIMI 48, Safety Analysis On-Treatmenta

<table>
<thead>
<tr>
<th>CrCL subgroup (mL/min)</th>
<th>Edoxaban 60 mg (N = 7,012)</th>
<th>Warfarin (N = 7,012)</th>
<th>HR (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>n</td>
<td>Number of Events</td>
<td>Event rate (%/year)</td>
</tr>
<tr>
<td>≥ 30 to ≤ 50</td>
<td>1,302</td>
<td>96</td>
<td>3.91</td>
</tr>
<tr>
<td>&gt; 50 to ≤ 70</td>
<td>2,093</td>
<td>148</td>
<td>3.31</td>
</tr>
<tr>
<td>&gt; 70 to ≤ 90</td>
<td>1,661</td>
<td>108</td>
<td>2.88</td>
</tr>
<tr>
<td>&gt; 90 to ≤ 110</td>
<td>927</td>
<td>29</td>
<td>1.33</td>
</tr>
<tr>
<td>&gt; 110 to ≤ 130</td>
<td>497</td>
<td>20</td>
<td>1.70</td>
</tr>
<tr>
<td>&gt; 130</td>
<td>462</td>
<td>13</td>
<td>1.18</td>
</tr>
</tbody>
</table>

Table 8: Number of Fatal Bleeding Events by creatinine clearance category in ENGAGE AF-TIMI 48, Safety Analysis On-Treatmenta

<table>
<thead>
<tr>
<th>CrCL subgroup (mL/min)</th>
<th>Edoxaban 60 mg (N = 7,012)</th>
<th>Warfarin (N = 7,012)</th>
<th>HR (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>n</td>
<td>Number of Events</td>
<td>Event rate (%/year)</td>
</tr>
<tr>
<td>≥ 30 to ≤ 50</td>
<td>1,302</td>
<td>9</td>
<td>0.36</td>
</tr>
<tr>
<td>&gt; 50 to ≤ 70</td>
<td>2,093</td>
<td>8</td>
<td>0.18</td>
</tr>
<tr>
<td>&gt; 70 to ≤ 90</td>
<td>1,661</td>
<td>10</td>
<td>0.26</td>
</tr>
<tr>
<td>&gt; 90 to ≤ 110</td>
<td>927</td>
<td>2</td>
<td>0.09</td>
</tr>
<tr>
<td>&gt; 110 to ≤ 130</td>
<td>497</td>
<td>1</td>
<td>0.08</td>
</tr>
<tr>
<td>&gt; 130</td>
<td>462</td>
<td>2</td>
<td>0.18</td>
</tr>
</tbody>
</table>

Table 9: Number of Intracranial Bleeding Events by creatinine clearance category in ENGAGE AF-TIMI 48, Safety Analysis On-Treatmenta

<table>
<thead>
<tr>
<th>CrCL subgroup (mL/min)</th>
<th>Edoxaban 60 mg (N = 7,012)</th>
<th>Warfarin (N = 7,012)</th>
<th>HR (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>n</td>
<td>Number of Events</td>
<td>Event rate (%/year)</td>
</tr>
<tr>
<td>≥ 30 to ≤ 50</td>
<td>1,302</td>
<td>16</td>
<td>0.64</td>
</tr>
<tr>
<td>&gt; 50 to ≤ 70</td>
<td>2,093</td>
<td>19</td>
<td>0.42</td>
</tr>
<tr>
<td>&gt; 70 to ≤ 90</td>
<td>1,661</td>
<td>17</td>
<td>0.44</td>
</tr>
<tr>
<td>&gt; 90 to ≤ 110</td>
<td>927</td>
<td>5</td>
<td>0.23</td>
</tr>
<tr>
<td>&gt; 110 to ≤ 130</td>
<td>497</td>
<td>2</td>
<td>0.17</td>
</tr>
<tr>
<td>&gt; 130</td>
<td>462</td>
<td>1</td>
<td>0.09</td>
</tr>
</tbody>
</table>

Abbreviations: N = number of subjects; mITT population overall study period; n = number of patients in subgroup
*HR not computed if number of events < 5 in one treatment group.
a On-Treatment: Time from first dose of study drug to last dose plus 3 days.
In subgroup analyses, for subjects in the 60 mg treatment group who were dose reduced to 30 mg in the ENGAGE AF-TIMI 48 study for body weight ≤ 60 kg, moderate renal impairment, or concomitant use of P-gp inhibitors, 104 (3.05% per year) of edoxaban 30 mg dose reduced subjects and 166 (4.85% per year) of warfarin dose reduced subjects had a major bleeding event [HR (95% CI): 0.63 (0.50, 0.81)].

In the ENGAGE AF-TIMI 48 study there was a significant improvement in Net Clinical Outcome (First Stroke, SEE, Major Bleed, or All-Cause Mortality; mITT population, overall study period) in favour of edoxaban, HR (95% CI): 0.89 (0.83, 0.96); p = 0.0024, when edoxaban 60 mg treatment group was compared to warfarin.

_Treatment of DVT, treatment of PE and the prevention of recurrent DVT and PE (VTE)_
The edoxaban clinical programme for VTE was designed to demonstrate the efficacy and safety of edoxaban in the treatment of DVT and PE, and the prevention of recurrent DVT and PE.

In the pivotal Hokusai-VTE study, 8,292 subjects were randomised to receive initial heparin therapy (enoxaparin or unfractionated heparin) followed by edoxaban 60 mg once daily or the comparator. In the comparator arm, subjects received initial heparin therapy concurrently with warfarin, titrated to a target INR of 2.0 to 3.0, followed by warfarin alone. The treatment duration was from 3 months up to 12 months, determined by the investigator based on the patient’s clinical features.

The majority of edoxaban treated patients were Caucasians (69.6%) and Asians (21.0%), 3.8% were Black, 5.3% were categorised as Other race.

The duration of therapy was at least 3 months for 3,718 (91.6%) edoxaban subjects versus 3,727 (91.4%) of warfarin subjects; at least 6 months for 3,495 (86.1%) of edoxaban subjects versus 3,491 (85.6%) of warfarin subjects; and 12 months for 1,643 (40.5%) edoxaban subjects versus 1,659 (40.4%) of warfarin subjects.

The primary efficacy endpoint was the recurrence of symptomatic VTE, defined as the composite of recurrent symptomatic DVT, non-fatal symptomatic PE and fatal PE in subjects during the 12-month study period. Secondary efficacy outcomes included the composite clinical outcome of recurrent VTE and all-cause mortality.

Edoxaban 30 mg once daily was used for subjects with one or more of the following clinical factors: moderate renal impairment (CrCL 30 - 50 mL/min); body weight ≤ 60 kg; concomitant use of specific P-gp inhibitors.

In the Hokusai-VTE study (Table 10) edoxaban was demonstrated to be non-inferior to warfarin for the primary efficacy outcome, recurrent VTE, which occurred in 130 of 4,118 subjects (3.2%) in the edoxaban group versus 146 of 4,122 subjects (3.5%) in the warfarin group [HR (95% CI): 0.89 (0.70, 1.13); p < 0.0001 for non-inferiority]. In the warfarin group, the median TTR (time in therapeutic range, INR 2.0 to 3.0) was 65.6%. For subjects presenting with PE (with or without DVT), 47 (2.8%) of edoxaban and 65 (3.9%) of warfarin subjects had a recurrent VTE [HR (95% CI): 0.73 (0.50, 1.06)].
Table 10: Efficacy Results from the Hokusai-VTE Study - mITT population, overall study period

<table>
<thead>
<tr>
<th>Primary endpoint(^a)</th>
<th>Edoxaban 60 mg (30 mg Dose Reduced) (N = 4,118)</th>
<th>Warfarin (N = 4,122)</th>
<th>Edoxaban vs Warfarin HR (95% CI)(^b) p-value(^c)</th>
</tr>
</thead>
<tbody>
<tr>
<td>All subjects with symptomatic recurrent VTE(^d), n (%)</td>
<td>130 (3.2)</td>
<td>146 (3.5)</td>
<td>0.89 (0.70, 1.13) p-value &lt; 0.0001 (non-inferiority)</td>
</tr>
<tr>
<td>PE with or without DVT</td>
<td>73 (1.8)</td>
<td>83 (2.0)</td>
<td></td>
</tr>
<tr>
<td>Fatal PE or Death where PE cannot be ruled out</td>
<td>24 (0.6)</td>
<td>24 (0.6)</td>
<td></td>
</tr>
<tr>
<td>Non-fatal PE</td>
<td>49 (1.2)</td>
<td>59 (1.4)</td>
<td></td>
</tr>
<tr>
<td>DVT only</td>
<td>57 (1.4)</td>
<td>63 (1.5)</td>
<td></td>
</tr>
</tbody>
</table>

Abbreviations: CI = Confidence Interval; DVT = deep vein thrombosis; mITT = modified intent-to-treat; HR = Hazard Ratio vs. warfarin; n = number of subjects with events; N = number of subjects in mITT population; PE = pulmonary embolism; VTE = venous thromboembolic events.

\(a\) The primary efficacy endpoint is adjudicated symptomatic recurrent VTE (i.e., the composite endpoint of DVT, non-fatal PE, and fatal PE).

\(b\) The HR, two-sided CI are based on the Cox proportional hazards regression model including treatment and the following randomisation stratification factors as covariates: presenting diagnosis (PE with or without DVT, DVT only), baseline risk factors (temporary factors, all others), and the need for 30 mg edoxaban/edoxaban placebo dose at randomisation (yes/no).

\(c\) The p-value is for the pre-defined non-inferiority margin of 1.5.

For the subjects who were dose reduced to 30 mg (predominantly low body weight or renal function) 15 (2.1%) edoxaban and 22 (3.1%) of warfarin subjects had a recurrent VTE [HR (95% CI): 0.69 (0.36, 1.34)].

The secondary composite endpoint of recurrent VTE and all-cause mortality occurred in 138 subjects (3.4%) in the edoxaban group and 158 subjects (3.9%) in the warfarin group [HR (95% CI): 0.87 (0.70, 1.10)].

The results for all-cause mortality (adjudicated deaths) in Hokusai-VTE were 136 (3.3%) for subjects taking edoxaban 60 mg (30 mg dose reduced) as opposed to 130 (3.2%) for warfarin.

In a pre-specified subgroup analysis of PE subjects 447 (30.6%) and 483 (32.2%) of edoxaban and warfarin treated subjects, respectively, were identified as having PE and NT-proBNP ≥ 500 pg/mL. The primary efficacy outcome occurred in 14 (3.1%) and 30 (6.2%) of edoxaban and warfarin subjects, respectively [HR (95% CI): 0.50 (0.26, 0.94)].

The efficacy results for pre-specified major subgroups (with dose reduction as required), including age, body weight, gender and status of renal function were consistent with the primary efficacy results for the overall population studied in the trial.

**Safety in patients with VTE (DVT and PE) in Hokusai-VTE**

The primary safety endpoint was clinically relevant bleeding (major or clinically relevant non-major).

Table 11 summarises adjudicated bleeding events for the safety analysis set on-treatment period.
There was a significant risk reduction in favour of edoxaban compared with warfarin for the primary safety endpoint of clinically relevant bleeding, a composite of major bleeding or clinically relevant non-major bleeding (CRNM), which occurred in 349 of 4,118 subjects (8.5%) in the edoxaban group and in 423 of 4,122 subjects (10.3%) in the warfarin group [HR (95% CI): 0.81 (0.71, 0.94); p = 0.004 for superiority].

Table 11: Bleeding Events in Hokusai-VTE Study - Safety Analysis On-Treatment Period

<table>
<thead>
<tr>
<th>Clinical Relevat Bleeding (Major and CRNM)b, n (%)</th>
<th>Edoxaban 60 mg (30 mg Dose Reduced) (N = 4,118)</th>
<th>Warfarin (N = 4,122)</th>
</tr>
</thead>
<tbody>
<tr>
<td>n</td>
<td>349 (8.5)</td>
<td>423 (10.3)</td>
</tr>
<tr>
<td>HR (95% CI)</td>
<td>0.81 (0.71, 0.94)</td>
<td></td>
</tr>
<tr>
<td>p-value</td>
<td>0.004 (for superiority)</td>
<td></td>
</tr>
</tbody>
</table>

**Major Bleeding** n (%)

| n                                                | 56 (1.4)                                    | 66 (1.6)            |
| HR (95% CI)                                       | 0.84 (0.59, 1.21)                           |                     |
| ICH fatal                                         | 0                                           | 6 (0.1)             |
| ICH non-fatal                                     | 5 (0.1)                                     | 12 (0.3)            |

**CRNM Bleeding**

| n                                                | 298 (7.2)                                   | 368 (8.9)           |
| HR (95% CI)                                       | 0.80 (0.68, 0.93)                           |                     |

**All Bleeding**

| n                                                | 895 (21.7)                                  | 1,056 (25.6)        |
| HR (95% CI)                                       | 0.82 (0.75, 0.90)                           |                     |

Abbreviations: ICH = Intracranial Haemorrhage, HR = Hazard Ratio vs. warfarin; CI = Confidence Interval; N = number of subjects in safety population; n = number of events; CRNM = clinically relevant non-major

a On-Treatment Period: Time from first dose of study drug to last dose plus 3 days.
b Primary Safety Endpoint: Clinically relevant bleeding (composite of major and clinically relevant non-major bleeding).

In subgroup analyses, for subjects who were dose reduced to 30 mg in the Hokusai-VTE study for body weight ≤ 60 kg, moderate renal impairment, or concomitant use of P-gp inhibitors, 58 (7.9%) of edoxaban 30 mg dose reduced subjects and 92 (12.8%) of warfarin subjects had a major bleeding or CRNM event [HR (95%): 0.62 (0.44, 0.86)].

In the Hokusai-VTE study the Net Clinical Outcome (Recurrent VTE, Major Bleed, or All-Cause Mortality; mITT population, overall study period) HR (95% CI) was 1.00 (0.85, 1.18) when edoxaban was compared to warfarin.

Patients undergoing cardioversion

A multicentre, prospective, randomised, open-label study with blinded endpoint evaluation (ENSURE-AF) was conducted which randomised 2199 subjects (oral anticoagulant naïve and pre-treated) with non-valvular atrial fibrillation scheduled for cardioversion, to compare edoxaban 60 mg once daily with enoxaparin/warfarin to maintain a therapeutic INR of 2.0 to 3.0 (randomised 1:1), mean TTR on warfarin was 70.8%. A total of 2149 subjects were treated with either edoxaban (N = 1067) or enoxaparin/warfarin (N = 1082). Subjects in the edoxaban treatment group received 30 mg once daily
if one or more of the following clinical factors were present: moderate renal impairment (CrCL 30 – 50 mL/min), low body weight (≤ 60 kg) or concomitant use of specific P-gp inhibitors. The majority of subjects in the edoxaban and warfarin groups had cardioversion performed (83.7% and 78.9%, respectively) or were auto-converted (6.6% and 8.6%, respectively). TEE-guided (within 3 days of initiation) or conventional cardioversion (at least 21 days of pre-treatment) was employed. Subjects were maintained on treatment for 28 days post cardioversion.

The primary efficacy outcome consisted of a composite of all stroke, SEE, MI and CV mortality. A total of 5 (0.5%, 95% CI 0.15% - 1.06%) events occurred in subjects in the edoxaban group (N = 1095) and 11 (1.0%, 95% CI 0.50% - 1.78%) events in the warfarin group (N = 1104); OR 0.46 (95% CI 0.12 - 1.43); ITT analysis set overall study period with mean duration of 66 days.

The primary safety outcome was a composite of major and CRNM bleeding. A total of 16 (1.5%, 95% CI 0.86% - 2.42%) events occurred in subjects in the edoxaban (N = 1067) group and 11 (1.0%, 95% CI 0.51% - 1.81%) events in the warfarin (N = 1082) group; OR 1.48 (95% CI 0.64 - 3.55); safety analysis set on-treatment period.

This exploratory study showed low rates of major and CRNM bleeding and thromboembolism in the two treatment groups in the setting of cardioversion.

**Paediatric population**

The European Medicines Agency has deferred the obligation to submit the results of studies with edoxaban in one or more subsets of the paediatric population in prevention of arterial thrombosis, treatment of thromboembolism and prevention of thromboembolism (see section 4.2 for information on paediatric use).

### 5.2 Pharmacokinetic properties

**Absorption**

Edoxaban is absorbed with peak plasma concentrations within 1 - 2 hours. The absolute bioavailability is approximately 62%. Food increases peak exposure to a varying extent, but has minimal effect on total exposure. Edoxaban was administered with or without food in the ENGAGE AF-TIMI 48 and the Hokusai-VTE studies. Edoxaban is poorly soluble at pH of 6.0 or higher. Co-administration of proton-pump inhibitors had no relevant impact on edoxaban exposure.

**Distribution**

Disposition is biphasic. The volume of distribution is 107 (19.9) L mean (SD). *In vitro* plasma protein binding is approximately 55%. There is no clinically relevant accumulation of edoxaban (accumulation ratio 1.14) with once daily dosing. Steady state concentrations are achieved within 3 days.

**Biotransformation**

Unchanged edoxaban is the predominant form in plasma. Edoxaban is metabolised via hydrolysis (mediated by carboxylesterase 1), conjugation or oxidation by CYP3A4/5 (< 10%). Edoxaban has three active metabolites, the predominant metabolite (M-4), formed by hydrolysis, is active and reaches less than 10% of the exposure of the parent compound in healthy subjects. Exposure to the other metabolites is less than 5%. Edoxaban is a substrate for the efflux transporter P-glycoprotein (P-gp), but not a substrate for uptake transporters such as organic anion transporter polypeptide OATP1B1, organic anion transporters OAT1 or OAT3 or organic cation transporter OCT2. Its active metabolite is a substrate for OATP1B1.

**Elimination**

In healthy subjects, the total clearance is estimated as 22 (± 3) L/hour; 50% is renally cleared (11 L/hour). Renal clearance accounts for approximately 35% of the administered dose. Metabolism and biliary/intestinal excretion account for the remaining clearance. The t½ for oral administration is 10 - 14 hours.
Linearity/non-linearity
Edoxaban displays approximately dose-proportional pharmacokinetics for doses of 15 mg to 60 mg in healthy subjects.

Special populations
Elderly
After taking renal function and body weight into account, age had no additional clinically significant effect on edoxaban pharmacokinetics in a population pharmacokinetic analysis of the pivotal Phase 3 study in NVAF (ENGAGE AF-TIMI 48).

Gender
After accounting for body weight, gender had no additional clinically significant effect on edoxaban pharmacokinetics in a population pharmacokinetic analysis of the Phase 3 study in NVAF (ENGAGE AF-TIMI 48).

Ethnic origin
In a population pharmacokinetic analysis of the ENGAGE AF-TIMI 48 study, peak and total exposure in Asian patients and non-Asian patients were comparable.

Renal impairment
The plasma AUCs for subjects with mild (CrCL > 50 - 80 mL/min), moderate (CrCL 30 - 50 mL/min) and severe (CrCL < 30 mL/min but not undergoing dialysis) renal impairment were increased by 32%, 74%, and 72%, respectively, relative to subjects with normal renal function. In patients with renal impairment the metabolite profile changes and a higher quantity of active metabolites are formed. There is a linear correlation between edoxaban plasma concentration and anti-FXa activity regardless of renal function.
Subjects with ESRD undergoing peritoneal dialysis had 93% higher total exposure compared with healthy subjects.
Population PK modeling indicates that exposure approximately doubles in patients with severe renal impairment (CrCL 15 – 29 mL/min) relative to patients with normal renal function.

Anti-FXa activity by CrCL category
Table 12 below shows the edoxaban anti-Factor Xa activity by CrCL category for each indication.
# Table 12: Edoxaban Anti-FXa activity by creatinine clearance

<table>
<thead>
<tr>
<th>Edoxaban Dose</th>
<th>CrCL (mL/min)</th>
<th>Edoxaban Anti-FXa activity post-dose (IU/mL)</th>
<th>Edoxaban Anti-FXa activity pre-dose (IU/mL)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Median [2.5 – 97.5% range]</td>
<td></td>
</tr>
<tr>
<td>Prevention of stroke and systemic embolism: NVAF</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>30 mg QD</td>
<td>≥ 30 to ≤ 50</td>
<td>2.92 [0.33 – 5.88]</td>
<td>0.53 [0.11 – 2.06]</td>
</tr>
<tr>
<td>60 mg QD*</td>
<td>&gt; 50 to ≤ 70</td>
<td>4.52 [0.38 – 7.64]</td>
<td>0.83 [0.16 – 2.61]</td>
</tr>
<tr>
<td></td>
<td>&gt; 70 to ≤ 90</td>
<td>4.12 [0.19 – 7.55]</td>
<td>0.68 [0.05 – 2.33]</td>
</tr>
<tr>
<td></td>
<td>&gt; 90 to ≤ 110</td>
<td>3.82 [0.36 – 7.39]</td>
<td>0.60 [0.14 – 3.57]</td>
</tr>
<tr>
<td></td>
<td>&gt; 110 to ≤ 130</td>
<td>3.16 [0.28 – 6.71]</td>
<td>0.41 [0.15 – 1.51]</td>
</tr>
<tr>
<td></td>
<td>&gt; 130</td>
<td>2.76 [0.12 – 6.10]</td>
<td>0.45 [0.00 – 3.10]</td>
</tr>
<tr>
<td>Treatment of DVT, treatment of PE and prevention of recurrent DVT and PE (VTE)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>30 mg QD</td>
<td>≥ 30 to ≤ 50</td>
<td>2.21 [0.14 – 4.47]</td>
<td>0.22 [0.00 – 1.09]</td>
</tr>
<tr>
<td>60 mg QD*</td>
<td>&gt; 50 to ≤ 70</td>
<td>3.42 [0.19 – 6.13]</td>
<td>0.34 [0.00 – 3.10]</td>
</tr>
<tr>
<td></td>
<td>&gt; 70 to ≤ 90</td>
<td>2.97 [0.24 – 5.82]</td>
<td>0.24 [0.00 – 1.77]</td>
</tr>
<tr>
<td></td>
<td>&gt; 90 to ≤ 110</td>
<td>2.82 [0.14 – 5.31]</td>
<td>0.20 [0.00 – 2.52]</td>
</tr>
<tr>
<td></td>
<td>&gt; 110 to ≤ 130</td>
<td>2.64 [0.13 – 5.57]</td>
<td>0.17 [0.00 – 1.86]</td>
</tr>
<tr>
<td></td>
<td>&gt; 130</td>
<td>2.39 [0.10 – 4.92]</td>
<td>0.13 [0.00 – 2.43]</td>
</tr>
</tbody>
</table>

*Dose reduction to 30 mg for low body weight ≤ 60 kg or specific concomitant P-glycoprotein (P-gp) inhibitors

1 Post-dose is equivalent to C<sub>max</sub> (post-dose samples were drawn 1 – 3 hours after edoxaban administration)

2 Pre-dose is equivalent to C<sub>min</sub>

Although treatment with edoxaban does not require routine monitoring, the effect on anticoagulation can be estimated by a calibrated quantitative anti-Factor Xa assay which may be useful in exceptional situations where knowledge of edoxaban exposure may help to inform clinical decisions, e.g. overdose and emergency surgery (see also section 4.4).

**Haemodialysis**

A 4 hour haemodialysis session reduced total edoxaban exposures by less than 9%.

**Hepatic impairment**

Patients with mild or moderate hepatic impairment exhibited comparable pharmacokinetics and pharmacodynamics to their matched healthy control group. Edoxaban has not been studied in patients with severe hepatic impairment (see section 4.2).
**Body weight**
In a population pharmacokinetic analysis of the ENGAGE AF-TIMI 48 study in NVAF, Cmax and AUC in patients with median low body weight (55 kg) were increased by 40% and 13%, respectively, as compared with patients with median high body weight (84 kg). In Phase 3 clinical studies (both NVAF and VTE indications) patients with body weight ≤ 60 kg had a 50% edoxaban dose reduction and had similar efficacy and less bleeding when compared to warfarin.

**Pharmacokinetic/pharmacodynamic relationship(s)**
PT, INR, aPTT and Anti-factor Xa correlate linearly with edoxaban concentrations.

### 5.3 Preclinical safety data

Non-clinical data reveal no special hazard for humans based on conventional studies of safety pharmacology, repeated dose toxicity, genotoxicity, carcinogenic potential, or phototoxicity.

**Reproductive toxicology**
Edoxaban showed vaginal haemorrhage at higher doses in rats and rabbits but had no effects in the reproductive performance of parent rats.

In rats, no effects on male or female fertility were seen.

In animal reproduction studies, rabbits showed increased incidence of gallbladder variations at a dosage of 200 mg/kg which is approximately 65 times the maximum recommended human dose (MRHD) of 60 mg/day based on total body surface area in mg/m². Increased post-implantation pregnancy losses occurred in rats at 300 mg/kg/day (approximately 49 times the MRHD) and in rabbits at 200 mg/kg/day (approximately 65 times the MRHD) respectively.

Edoxaban was excreted in the breast milk of lactating rats.

**Environmental Risk Assessment (ERA)**
The active substance edoxaban tosilate is persistent in the environment (for instructions on disposal see section 6.6).

### 6. PHARMACEUTICAL PARTICULARS

#### 6.1 List of excipients

**Tablet core:**
- Mannitol (E421)
- Pregelatinised starch
- Crospovidone
- Hydroxypropylcellulose
- Magnesium stearate (E470b)

**Film-coat:**
- Hypromellose (E464)
- Macrogol 8000
- Titanium dioxide (E171)
- Talc
- Carnauba wax
- Iron oxide yellow (E172)

#### 6.2 Incompatibilities

Not applicable.
6.3 Shelf life

5 years

6.4 Special precautions for storage

This medicinal product does not require any special storage conditions.

6.5 Nature and contents of container

PVC/Aluminium blisters. Cartons of 10, 14, 28, 30, 56, 60, 84, 90, 98, 100 film-coated tablets. PVC/Aluminium perforated unit dose blisters of 10 x 1, 50 x 1 and 100 x 1 film-coated tablets.

Not all pack sizes may be marketed.

6.6 Special precautions for disposal

Any unused medicinal product or waste material should be disposed of in accordance with local requirements.

7. MARKETING AUTHORITY

Daiichi Sankyo Europe GmbH
Zielstattstrasse 48
81379 Munich
Germany

8. MARKETING AUTHORIZATION NUMBER(S)

EU/1/15/993/003
EU/1/15/993/017-028

9. DATE OF FIRST AUTHORIZATION/RENEWAL OF THE AUTHORIZATION

Date of first authorisation: 19 June 2015

10. DATE OF REVISION OF THE TEXT

Detailed information on this medicinal product is available on the website of the European Medicines Agency http://www.ema.europa.eu.
ANNEX II

A. MANUFACTURER(S) RESPONSIBLE FOR BATCH RELEASE

B. CONDITIONS OR RESTRICTIONS REGARDING SUPPLY AND USE

C. OTHER CONDITIONS AND REQUIREMENTS OF THE MARKETING AUTHORISATION

D. CONDITIONS OR RESTRICTIONS WITH REGARD TO THE SAFE AND EFFECTIVE USE OF THE MEDICINAL PRODUCT
A. MANUFACTURER RESPONSIBLE FOR BATCH RELEASE

Name and address of the manufacturer responsible for batch release

Daiichi Sankyo Europe GmbH
Luitpoldstrasse 1
85276 Pfaffenhofen, Bayern
Germany

B. CONDITIONS OR RESTRICTIONS REGARDING SUPPLY AND USE

Medicinal product subject to medical prescription.

C. OTHER CONDITIONS AND REQUIREMENTS OF THE MARKETING AUTHORIZATION

- Periodic safety update reports

The requirements for submission of periodic safety update reports for this medicinal product are set out in the list of Union reference dates (EURD list) provided for under Article 107c(7) of Directive 2001/83/EC and any subsequent updates published on the European medicines web-portal.

D. CONDITIONS OR RESTRICTIONS WITH REGARD TO THE SAFE AND EFFECTIVE USE OF THE MEDICINAL PRODUCT

- Risk Management Plan (RMP)

The MAH shall perform the required pharmacovigilance activities and interventions detailed in the agreed RMP presented in Module 1.8.2 of the Marketing Authorisation and any agreed subsequent updates of the RMP.

An updated RMP should be submitted:
- At the request of the European Medicines Agency;
- Whenever the risk management system is modified, especially as the result of new information being received that may lead to a significant change to the benefit/risk profile or as the result of an important (pharmacovigilance or risk minimisation) milestone being reached.
• **Additional risk minimisation measures**

Prior to launch of Lixiana in each Member State, the Marketing Authorisation Holder (MAH) must agree the content and format of the educational programme, including communication media, distribution modalities, and any other aspects of the programme, with the National Competent Authority.

The educational programme is aimed at mitigating the risk of serious bleeds or haemorrhage in patients treated with Lixiana by ensuring prescriber awareness and providing guidance on appropriate patient selection, correct dosing as well as management of the risk.

The programme is also aimed at ensuring that the healthcare professionals who intend to prescribe Lixiana are aware of the Patient alert card and that the card is to be given to and reviewed with all patients treated with Lixiana.

The MAH shall ensure that in each Member State where Lixiana is marketed, all healthcare professionals who are expected to use Lixiana are provided with the following educational material:

- The Summary of Product Characteristics
- Prescriber Guide for healthcare professionals
- Patient alert card

The Prescriber Guide for healthcare professionals shall contain the following key elements:

- Relevant information on the risk of bleeding
- Details of the population potentially at higher risk of bleeding
- Contraindications
- Recommendations for dose adjustment in at risk populations, including patients with renal or hepatic impairment, low body weight and concomitant use of some P-gp inhibitors
- Guidance on switching from or to Lixiana treatment
- Guidance regarding surgery or invasive procedure, and temporary discontinuation
- Management of overdose situations and haemorrhage
- Use of coagulation tests and their interpretation
- That all patients should be provided with a Patient alert card and be counselled about:
  - The signs or symptoms of bleeding and when to seek attention from a healthcare provider
  - Importance of treatment compliance
  - Necessity to carry the Patient alert card with them at all times
  - The need to inform Health Care Professionals that they are taking Lixiana if they need to have any surgery or invasive procedure

The Patient alert card should contain the following key safety messages:

- The signs or symptoms of bleeding and when to seek attention
- Importance of treatment compliance
- Necessity to carry the Patient alert card with them at all times
- The need to inform Health Care Professionals that they are taking Lixiana if they need to have any surgery or invasive procedure
A. LABELLING
**PARTICULARS TO APPEAR ON THE OUTER PACKAGING**

**OUTER CARTON FOR 15 MG**

1. **NAME OF THE MEDICINAL PRODUCT**
   Lixiana 15 mg film-coated tablets
edoxaban

2. **STATEMENT OF ACTIVE SUBSTANCE(S)**
   Each film-coated tablet contains 15 mg edoxaban (as tosilate).

3. **LIST OF EXCIPIENTS**

4. **PHARMACEUTICAL FORM AND CONTENTS**
   10 film-coated tablets
   10 x 1 film-coated tablets

5. **METHOD AND ROUTE(S) OF ADMINISTRATION**
   Read the package leaflet before use.
   Oral use.

6. **SPECIAL WARNING THAT THE MEDICINAL PRODUCT MUST BE STORED OUT OF THE SIGHT AND REACH OF CHILDREN**
   Keep out of the sight and reach of children.

7. **OTHER SPECIAL WARNING(S), IF NECESSARY**

8. **EXPIRY DATE**
   EXP:

9. **SPECIAL STORAGE CONDITIONS**

10. **SPECIAL PRECAUTIONS FOR DISPOSAL OF UNUSED MEDICINAL PRODUCTS OR WASTE MATERIALS DERIVED FROM SUCH MEDICINAL PRODUCTS, IF APPROPRIATE**
11. NAME AND ADDRESS OF THE MARKETING AUTHORISATION HOLDER

Daiichi Sankyo Europe GmbH
81366 Munich
Germany

12. MARKETING AUTHORISATION NUMBER(S)

EU/1/15/993/001 10 film-coated tablets
EU/1/15/993/016 10 x 1 film-coated tablets

13. BATCH NUMBER

Lot:

14. GENERAL CLASSIFICATION FOR SUPPLY

15. INSTRUCTIONS ON USE

16. INFORMATION IN BRAILLE

Lixiana 15 mg

17. UNIQUE IDENTIFIER – 2D BARCODE

2D barcode carrying the unique identifier included.

18. UNIQUE IDENTIFIER - HUMAN READABLE DATA

PC:
SN:
NN:

A QR Code and the following text may be printed on the inner lid of the carton:

Lixiana should be taken once daily and ideally at the same time every day
Take as recommended by your doctor:

☐ Either in the morning    ☐ Or at noon    ☐ Or in the evening

For more information please visit:

www.dspatient.eu

Or scan the QR code with your smartphone:

QR code to be included

These are your last remaining tablets.
Please consult your doctor for a new prescription of Lixiana.
Do not stop taking Lixiana unless advised by your doctor.
### 1. NAME OF THE MEDICINAL PRODUCT

Lixiana 15 mg film-coated tablets
edoxaban

### 2. NAME OF THE MARKETING AUTHORISATION HOLDER

Daiichi-Sankyo (logo)

### 3. EXPIRY DATE

EXP:

### 4. BATCH NUMBER

Lot:

### 5. OTHER
<table>
<thead>
<tr>
<th>MINIMUM PARTICULARS TO APPEAR ON BLISTERS OR STRIPS</th>
</tr>
</thead>
<tbody>
<tr>
<td>UNIT DOSE BLISTER (10 x 1 TABLETS) FOR 15 MG</td>
</tr>
</tbody>
</table>

1. **NAME OF THE MEDICINAL PRODUCT**

Lixiana 15 mg film-coated tablets
edoxaban

2. **NAME OF THE MARKETING AUTHORISATION HOLDER**

Daiichi-Sankyo (logo)

3. **EXPIRY DATE**

EXP:

4. **BATCH NUMBER**

Lot:

5. **OTHER**
**PARTICULARS TO APPEAR ON THE OUTER PACKAGING**

**OUTER CARTON FOR 30 MG**

1. **NAME OF THE MEDICINAL PRODUCT**

Lixiana 30 mg film-coated tablets
edoxaban

2. **STATEMENT OF ACTIVE SUBSTANCE(S)**

Each film-coated tablet contains 30 mg edoxaban (as tosilate).

3. **LIST OF EXCIPIENTS**

4. **PHARMACEUTICAL FORM AND CONTENTS**

<table>
<thead>
<tr>
<th>Number of Tablets</th>
</tr>
</thead>
<tbody>
<tr>
<td>10</td>
</tr>
<tr>
<td>14</td>
</tr>
<tr>
<td>28</td>
</tr>
<tr>
<td>30</td>
</tr>
<tr>
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<td>98</td>
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<tr>
<td>100</td>
</tr>
<tr>
<td>10 x 1</td>
</tr>
<tr>
<td>50 x 1</td>
</tr>
<tr>
<td>100 x 1</td>
</tr>
</tbody>
</table>

5. **METHOD AND ROUTE(S) OF ADMINISTRATION**

Read the package leaflet before use.

Oral use.

6. **SPECIAL WARNING THAT THE MEDICINAL PRODUCT MUST BE STORED OUT OF THE SIGHT AND REACH OF CHILDREN**

Keep out of the sight and reach of children.

7. **OTHER SPECIAL WARNING(S), IF NECESSARY**

8. **EXPIRY DATE**

EXP:
9. SPECIAL STORAGE CONDITIONS

10. SPECIAL PRECAUTIONS FOR DISPOSAL OF UNUSED MEDICINAL PRODUCTS OR WASTE MATERIALS DERIVED FROM SUCH MEDICINAL PRODUCTS, IF APPROPRIATE

11. NAME AND ADDRESS OF THE MARKETING AUTHORISATION HOLDER

Daiichi Sankyo Europe GmbH
81366 Munich
Germany

12. MARKETING AUTHORISATION NUMBER(S)

- EU/1/15/993/002 10 film-coated tablets
- EU/1/15/993/004 14 film-coated tablets
- EU/1/15/993/005 28 film-coated tablets
- EU/1/15/993/006 30 film-coated tablets
- EU/1/15/993/007 56 film-coated tablets
- EU/1/15/993/008 60 film-coated tablets
- EU/1/15/993/009 84 film-coated tablets
- EU/1/15/993/010 90 film-coated tablets
- EU/1/15/993/011 98 film-coated tablets
- EU/1/15/993/012 100 film-coated tablets
- EU/1/15/993/013 10 x 1 film-coated tablets
- EU/1/15/993/014 50 x 1 film-coated tablets
- EU/1/15/993/015 100 x 1 film-coated tablets

13. BATCH NUMBER

Lot:

14. GENERAL CLASSIFICATION FOR SUPPLY

15. INSTRUCTIONS ON USE

16. INFORMATION IN BRAILLE

Lixiana 30 mg

17. UNIQUE IDENTIFIER – 2D BARCODE

2D barcode carrying the unique identifier included.
A QR Code and the following text may be printed on the inner lid of the carton:

Lixiana should be taken once daily and ideally at the same time every day

Take as recommended by your doctor:

☐ Either in the morning  ☐ Or at noon  ☐ Or in the evening

For more information please visit:

www.dspatient.eu

Or scan the QR code with your smartphone:

QR code to be included

These are your last remaining tablets. Please consult your doctor for a new prescription of Lixiana. Do not stop taking Lixiana unless advised by your doctor.
MINIMUM PARTICULARS TO APPEAR ON BLISTERS OR STRIPS

BLISTER OF 10 FILM-COATED TABLETS FOR 30 MG

1. NAME OF THE MEDICINAL PRODUCT

Lixiana 30 mg film-coated tablets
edoxaban

2. NAME OF THE MARKETING AUTHORISATION HOLDER

Daiichi-Sankyo (logo)

3. EXPIRY DATE

EXP:

4. BATCH NUMBER

Lot:

5. OTHER
<table>
<thead>
<tr>
<th>MINIMUM PARTICULARS TO APPEAR ON BLISTERS OR STRIPS</th>
</tr>
</thead>
<tbody>
<tr>
<td>BLISTER OF 14 FILM-COATED TABLETS FOR 30 MG</td>
</tr>
</tbody>
</table>

1. **NAME OF THE MEDICINAL PRODUCT**

Lixiana 30 mg film-coated tablets
edoxaban

2. **NAME OF THE MARKETING AUTHORISATION HOLDER**

Daiichi-Sankyo (logo)

3. **EXPIRY DATE**

EXP:

4. **BATCH NUMBER**

Lot:

5. **OTHER**

MON, TUE, WED, THU, FRI, SAT, SUN
<table>
<thead>
<tr>
<th>MINIMUM PARTICULARS TO APPEAR ON BLISTERS OR STRIPS</th>
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<tr>
<td>UNIT DOSE BLISTER (10 x 1 TABLETS) FOR 30 MG</td>
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1. **NAME OF THE MEDICINAL PRODUCT**

Lixiana 30 mg film-coated tablets
edoxaban

2. **NAME OF THE MARKETING AUTHORIZATION HOLDER**

Daiichi-Sankyo (logo)

3. **EXPIRY DATE**

EXP:

4. **BATCH NUMBER**

Lot:

5. **OTHER**
PARTICULARS TO APPEAR ON THE OUTER PACKAGING

OUTER CARTON FOR 60 MG

1. NAME OF THE MEDICINAL PRODUCT

Lixiana 60 mg film-coated tablets
edoxaban

2. STATEMENT OF ACTIVE SUBSTANCE(S)

Each film-coated tablet contains 60 mg edoxaban (as tosilate).

3. LIST OF EXCIPIENTS

4. PHARMACEUTICAL FORM AND CONTENTS

10 film-coated tablets
14 film-coated tablets
28 film-coated tablets
30 film-coated tablets
56 film-coated tablets
60 film-coated tablets
84 film-coated tablets
90 film-coated tablets
98 film-coated tablets
100 film-coated tablets
10 x 1 film-coated tablets
50 x 1 film-coated tablets
100 x 1 film-coated tablets

5. METHOD AND ROUTE(S) OF ADMINISTRATION

Read the package leaflet before use.
Oral use.

6. SPECIAL WARNING THAT THE MEDICINAL PRODUCT MUST BE STORED OUT OF THE SIGHT AND REACH OF CHILDREN

Keep out of the sight and reach of children.

7. OTHER SPECIAL WARNING(S), IF NECESSARY

8. EXPIRY DATE

EXP:
9. **SPECIAL STORAGE CONDITIONS**

10. **SPECIAL PRECAUTIONS FOR DISPOSAL OF UNUSED MEDICINAL PRODUCTS OR WASTE MATERIALS DERIVED FROM SUCH MEDICINAL PRODUCTS, IF APPROPRIATE**

11. **NAME AND ADDRESS OF THE MARKETING AUTHORISATION HOLDER**

Daiichi Sankyo Europe GmbH  
81366 Munich  
Germany

12. **MARKETING AUTHORISATION NUMBER(S)**

<table>
<thead>
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<tr>
<td>EU/1/15/993/003</td>
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<td>50 x 1 film-coated tablets</td>
</tr>
<tr>
<td>EU/1/15/993/028</td>
<td>100 x1 film-coated tablets</td>
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13. **BATCH NUMBER**

Lot:

14. **GENERAL CLASSIFICATION FOR SUPPLY**

15. **INSTRUCTIONS ON USE**

16. **INFORMATION IN BRAILLE**

Lixiana 60 mg

17. **UNIQUE IDENTIFIER – 2D BARCODE**

2D barcode carrying the unique identifier included.
A QR Code and the following text may be printed on the inner lid of the carton:

Lixiana should be taken once daily and ideally at the same time every day

Take as recommended by your doctor:

☐ Either in the morning ☐ Or at noon ☐ Or in the evening

For more information please visit:

www.dspatient.eu

Or scan the QR code with your smartphone:

QR code to be included

These are your last remaining tablets.
Please consult your doctor for a new prescription of Lixiana.
Do not stop taking Lixiana unless advised by your doctor.
<table>
<thead>
<tr>
<th>MINIMUM PARTICULARS TO APPEAR ON BLISTERS OR STRIPS</th>
</tr>
</thead>
<tbody>
<tr>
<td>BLISTER OF 10 FILM-COATED TABLETS FOR 60 MG</td>
</tr>
</tbody>
</table>

1. **NAME OF THE MEDICINAL PRODUCT**

   Lixiana 60 mg film-coated tablets  
edoxaban

2. **NAME OF THE MARKETING AUTHORISATION HOLDER**

   Daiichi-Sankyo (logo)

3. **EXPIRY DATE**

   EXP:

4. **BATCH NUMBER**

   Lot:

5. **OTHER**
<table>
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<tr>
<th>MINIMUM PARTICULARS TO APPEAR ON BLISTERS OR STRIPS</th>
</tr>
</thead>
<tbody>
<tr>
<td>BLISTER OF 14 FILM-COATED TABLETS FOR 60 MG</td>
</tr>
</tbody>
</table>

1. **NAME OF THE MEDICINAL PRODUCT**

Lixiana 60 mg film-coated tablets
edoxaban

2. **NAME OF THE MARKETING AUTHORISATION HOLDER**

Daiichi-Sankyo (logo)

3. **EXPIRY DATE**

EXP:

4. **BATCH NUMBER**

Lot:

5. **OTHER**

MON, TUE, WED, THU, FRI, SAT, SUN
<table>
<thead>
<tr>
<th>MINIMUM PARTICULARS TO APPEAR ON BLISTERS OR STRIPS</th>
</tr>
</thead>
<tbody>
<tr>
<td>UNIT DOSE BLISTER (10 x 1 TABLETS) FOR 60 MG</td>
</tr>
</tbody>
</table>

1. **NAME OF THE MEDICINAL PRODUCT**

Lixiana 60 mg film-coated tablets
edoxaban

2. **NAME OF THE MARKETING AUTHORISATION HOLDER**

Daiichi-Sankyo (logo)

3. **EXPIRY DATE**

EXP:

4. **BATCH NUMBER**

Lot:

5. **OTHER**
PATIENT ALERT CARD

**Lixiana**
film-coated tablets
**edoxaban**

Please keep this card with you at all times.

Present it to your healthcare professional, pharmacist, surgeon or dentist before any medical treatment or intervention.

PATIENT INFORMATION

Patient name:
Date of birth:
In case of emergency, please contact:
Name:
Phone no:

TREATMENT INFORMATION
(To be completed by physician)

Lixiana has been prescribed at a once-daily dose of: mg

Started on: / (mm/yy)
Blood type:
Other medications/conditions:

PRESCRIBER INFORMATION

For more information or in case of emergency, please contact:

Physician’s name:
Phone number, practice stamp:
Signature of physician:

INFORMATION FOR HEALTHCARE PROFESSIONALS

- Lixiana is an oral anticoagulant factor Xa inhibitor.
- When an invasive procedure is required, Lixiana should be stopped at least 24 hours beforehand, and appropriate caution exercised.
• Lixiana may increase the risk of bleeding. In case of clinically significant bleeding, stop treatment immediately.
• Coagulation tests such as INR, prothrombin time (PT), or activated partial thromboplastin time (aPTT) are not a useful measure of the effect of Lixiana. However, a calibrated anti-Factor Xa assay may help inform clinical decisions.

Please consult the Summary of Product Characteristics (SmPC) for more information.

ABOUT YOUR TREATMENT

You have been prescribed Lixiana, an anticoagulant medicine, which thins your blood and helps prevent you from suffering blood clots. It is important that you take your medication exactly as instructed by your doctor.

• If you miss a dose, take it immediately and then continue the following day as normal – do not take double the prescribed dose in a single day.
• Do not start any other medications (including over the counter) without consulting your doctor.
• Do not stop taking Lixiana without consulting your doctor as this can increase your risk of developing a blood clot.
• Please read the Patient Information Leaflet found inside each pack of Lixiana.

WHEN TO SEEK MEDICAL ADVICE

RISK OF BLEEDING

Taking an anticoagulant medicine such as Lixiana can increase your risk of bleeding. It is therefore important to be aware of the possible signs and symptoms of bleeding and to speak to your doctor immediately if you experience any of the following:

• Bruising or bleeding under the skin
• Blood in the urine
• Coughing up blood
• Vomiting blood or material that looks like ground coffee
• Nose bleeds or cuts that take a long time to stop bleeding
• Tar-coloured stools
• Dizziness or sudden headache
• Unexplained tiredness
• Abnormal vaginal bleeding, including heavier or prolonged menses

Please talk to your doctor if you experience any unusual symptoms.
B. PACKAGE LEAFLET
This medicine is subject to additional monitoring. This will allow quick identification of new safety information. You can help by reporting any side effects you may get. See the end of section 4 for how to report side effects.

Read all of this leaflet carefully before you start taking this medicine because it contains important information for you.
- Keep this leaflet. You may need to read it again.
- If you have any further questions, ask your doctor or pharmacist.
- This medicine has been prescribed for you only. Do not pass it on to others. It may harm them, even if their signs of illness are the same as yours.
- If you get any side effects, talk to your doctor or pharmacist. This includes any possible side effects not listed in this leaflet. See section 4.

What is in this leaflet
1. What Lixiana is and what it is used for
2. What you need to know before you take Lixiana
3. How to take Lixiana
4. Possible side effects
5. How to store Lixiana
6. Contents of the pack and other information

1. What Lixiana is and what it is used for

Lixiana contains the active substance edoxaban and belongs to a group of medicines called anticoagulants. This medicine helps to prevent blood clots from forming. It works by blocking the activity of factor Xa, which is an important component of blood clotting.

Lixiana is used in adults to:

- **prevent blood clots in the brain** (stroke) and **other blood vessels in the body** if you have a form of irregular heart rhythm called nonvalvular atrial fibrillation and at least one additional risk factor.

- **treat blood clots in the veins of the legs** (deep vein thrombosis) and **in the blood vessels in the lungs** (pulmonary embolism), and to **prevent blood clots from re-occurring** in the blood vessels in the legs and/or lungs.
2. What you need to know before you take Lixiana

Do not take Lixiana:

- if you are allergic to edoxaban or any of the other ingredients of this medicine (listed in section 6)
- if you are actively bleeding
- if you have a disease or condition that increases the risk of serious bleeding (e.g. stomach ulcer, injury or bleeding in the brain, or recent surgery of the brain or eyes)
- if you are taking other medicines to prevent blood clotting (e.g. warfarin, dabigatran, rivaroxaban, apixaban or heparin), except when changing anticoagulant treatment or while getting heparin through a venous or arterial line to keep it open
- if you have a liver disease which leads to an increased risk of bleeding
- if you have uncontrolled high blood pressure
- if you are pregnant or breast feeding

Warnings and precautions

Talk to your doctor or pharmacist before taking Lixiana,

- if you have an increased risk of bleeding, as could be the case if you have any of the following conditions:
  - endstage kidney disease or if you are on dialysis
  - severe liver disease
  - bleeding disorders
  - a problem with the blood vessels in the back of your eyes (retinopathy)
  - recent bleeding in your brain (intracranial or intracerebral bleeding)
  - problems with the blood vessels in your brain or spinal column
- if you have a mechanical heart valve

Lixiana 15 mg is only to be used when changing from Lixiana 30 mg to a vitamin K antagonist (e.g. warfarin) (see section 3. How to take Lixiana).

If you need to have an operation:
It is very important to take Lixiana before and after the operation exactly at the times you have been told by your doctor. If possible, Lixiana should be stopped at least 24 hours before an operation. Your doctor will determine when to restart Lixiana.

Children and adolescents
Lixiana is not recommended in children and adolescents under 18 years of age. There is no information on its use in children and adolescents.

Other medicines and Lixiana
Tell your doctor or pharmacist if you are taking, have recently taken or might take any other medicines.

If you are taking any of the following:
- some medicines for fungal infections (e.g. ketoconazole)
- medicines to treat abnormal heart beat (e.g. dronedarone, quinidine, verapamil)
- other medicines to reduce blood clotting (e.g. heparin, clopidogrel or vitamin K antagonists such as warfarin, acenocoumarol, phenprocoumon or dabigatran, rivaroxaban, apixaban)
- antibiotic medicines (e.g. erythromycin)
- medicines to prevent organ rejection after transplantation (e.g. ciclosporin)
- anti-inflammatory and pain-relieving medicines (e.g. naproxen or acetylsalicylic acid (aspirin))

Tell your doctor before taking Lixiana, because these medicines may increase the effects of Lixiana and the chance of unwanted bleeding. Your doctor will decide, if you should be treated with Lixiana and if you should be kept under observation.
If you are taking any of the following:
- some medicines for treatment of epilepsy (e.g. phenytoin, carbamazepine, phenobarbital)
- St John’s Wort, a herbal product used for anxiety and mild depression
- rifampicin, an antibiotic

**If any of the above apply to you, tell your doctor** before taking Lixiana, because the effect of Lixiana may be reduced. Your doctor will decide if you should be treated with Lixiana and if you should be kept under observation.

**Pregnancy and breast-feeding**
Do not take Lixiana if you are pregnant or breast-feeding. If there is a chance that you could become pregnant, use a reliable contraceptive while you are taking Lixiana. If you become pregnant while you are taking Lixiana, immediately tell your doctor, who will decide how you should be treated.

**Driving and using machines**
Lixiana has no or negligible effects on your ability to drive or use machines.

### 3. How to take Lixiana

Always take this medicine exactly as your doctor or pharmacist has told you. Check with your doctor or pharmacist if you are not sure.

**How much to take**
The recommended dose is one **60 mg** tablet once daily.

- **If you have impaired kidney function**, the dose may be reduced to one **30 mg** tablet once daily by your doctor.
- **If your body weight is 60 kg or lower**, the recommended dose is one **30 mg** tablet once daily.
- **If your doctor has prescribed medicines known as P-gp inhibitors**: ciclosporin, dronedarone, erythromycin, or ketoconazole, the recommended dose is one **30 mg** tablet once daily.

**How to take the tablet**
Swallow the tablet, preferably with water. Lixiana can be taken with or without food.

**Your doctor may change your anticoagulant treatment as follows:**

*Changing from vitamin K antagonists (e.g. warfarin) to Lixiana*
Stop taking the vitamin K antagonist (e.g. warfarin). Your doctor will need to do blood measurements and will instruct you when to start taking Lixiana.

*Changing from non-VKA oral anticoagulants (dabigatran, rivaroxaban, or apixaban) to Lixiana*
Stop taking the previous medicines (e.g. dabigatran, rivaroxaban, or apixaban) and start Lixiana at the time of the next scheduled dose.

*Changing from parenteral anticoagulants (e.g. heparin) to Lixiana*
Stop taking the anticoagulant (e.g. heparin) and start Lixiana at the time of the next scheduled anticoagulant dose.

*Changing from Lixiana to vitamin K antagonists (e.g. warfarin)*
If you currently take 60 mg Lixiana:
Your doctor will tell you to reduce your dose of Lixiana to a 30 mg tablet once daily and to take it together with a vitamin K antagonist (e.g. warfarin). Your doctor will need to do blood measurements and will instruct you when to stop taking Lixiana.
If you currently take 30 mg (dose reduced) Lixiana:
Your doctor will tell you to reduce your dose of Lixiana to a 15 mg tablet once daily and to take it together with a vitamin K antagonist (e.g. warfarin). Your doctor will need to do blood measurements and will instruct you when to stop taking Lixiana.

Changing from Lixiana to non-VKA oral anticoagulants (dabigatran, rivaroxaban, or apixaban)
Stop taking Lixiana and start the non-VKA anticoagulant (e.g. dabigatran, rivaroxaban, or apixaban) at the time of the next scheduled dose of Lixiana.

Changing from Lixiana to parenteral anticoagulants (e.g. heparin)
Stop taking Lixiana and start the parenteral anticoagulant (e.g. heparin) at the time of the next scheduled dose of Lixiana.

Patients undergoing cardioversion:
If your abnormal heartbeat needs to be restored to normal by a procedure called cardioversion, take Lixiana at the times your doctor tells you to prevent blood clots in the brain and other blood vessels in your body.

If you take more Lixiana than you should
Tell your doctor immediately if you have taken too many Lixiana tablets.
If you take more Lixiana than recommended, you may have an increased risk of bleeding.

If you forget to take Lixiana
You should take the tablet immediately and then continue the following day with the once daily tablet as usual. Do not take a double dose on the same day to make up for a forgotten dose.

If you stop taking Lixiana
Do not stop taking Lixiana without talking to your doctor first, because Lixiana treats and prevents serious conditions.

If you have any further questions on the use of this medicine, ask your doctor or pharmacist.

4. Possible side effects

Like all medicines, this medicine can cause side effects, although not everybody gets them.

Like other similar medicines (medicines to reduce blood clotting), Lixiana may cause bleeding which may potentially be life-threatening. In some cases the bleeding may not be obvious.

If you experience any bleeding event that does not stop by itself or if you experience signs of excessive bleeding (exceptional weakness, tiredness, paleness, dizziness, headache or unexplained swelling) consult your doctor immediately.
Your doctor may decide to keep you under closer observation or change your medicine.

Overall list of possible side effects:

Common (may affect up to 1 in 10 people):
- Stomach ache
- Abnormal liver blood tests
- Bleeding from the skin or under the skin
- Anaemia (low levels of red blood cells)
- Bleeding from the nose
- Bleeding from the vagina
- Rash
• Bleeding in the bowel
• Bleeding from the mouth and/or throat
• Blood found in your urine
• Bleeding following an injury (puncture)
• Bleeding in the stomach
• Dizziness
• Feeling sick
• Headache
• Itching

**Uncommon** (may affect up to 1 in 100 people):
• Other types of bleeding
• Bleeding in the eyes
• Bleeding from a surgical wound following an operation
• Blood in the spit when coughing
• Bleeding in the brain
• Reduced number of platelets in your blood (which can affect clotting)
• Allergic reaction
• Hives

**Rare** (may affect up to 1 in 1,000 people):
• Bleeding in the muscles
• Bleeding in joints
• Bleeding in the abdomen
• Bleeding in the heart
• Bleeding inside the skull
• Bleeding following a surgical procedure

**Reporting of side effects**
If you get any side effects, talk to your doctor or pharmacist. This includes any possible side effects not listed in this leaflet. You can also report side effects directly via [the national reporting system listed in Appendix V](#). By reporting side effects you can help provide more information on the safety of this medicine.

5. **How to store Lixiana**

Keep this medicine out of the sight and reach of children.

Do not use this medicine after the expiry date which is stated on the carton and on each blister after EXP. The expiry date refers to the last day of that month.

This medicine does not require any special storage conditions.

Do not throw away any medicines via wastewater or household waste. Ask your pharmacist how to throw away medicines you no longer use. These measures will help protect the environment.

6. **Contents of the pack and other information**

**What Lixiana contains**
- The active substance is edoxaban (as tosilate).

  *Lixiana 15 mg:* Each tablet contains 15 mg edoxaban (as tosilate).
  *Lixiana 30 mg:* Each tablet contains 30 mg edoxaban (as tosilate).
Lixiana 60 mg: Each tablet contains 60 mg edoxaban (as tosilate).

The other ingredients are:

Lixiana 15 mg: Tablet core: mannitol (E421), pregelatinised starch, crospovidone, hydroxypropylcellulose, magnesium stearate (E470b).
Lixiana 30 mg: Tablet core: mannitol (E421), pregelatinised starch, crospovidone, hydroxypropylcellulose, magnesium stearate (E470b).
Lixiana 60 mg: Tablet core: mannitol (E421), pregelatinised starch, crospovidone, hydroxypropylcellulose, magnesium stearate (E470b).

Film coat:

Lixiana 15 mg: hypromellose (E464), macrogol 8000, titanium dioxide (E171), talc, carnauba wax, iron oxide red (E172), iron oxide yellow (E172).
Lixiana 30 mg: hypromellose (E464), macrogol 8000, titanium dioxide (E171), talc, carnauba wax, iron oxide red (E172).
Lixiana 60 mg: hypromellose (E464), macrogol 8000, titanium dioxide (E171), talc, carnauba wax, iron oxide yellow (E172).

What Lixiana looks like and contents of the pack

Lixiana 15 mg film-coated tablets are orange, round-shaped (6.7 mm diameter) and debossed with “DSC L15” on one side.
They come in blisters in cartons of 10 film-coated tablets or unit dose blisters in cartons of 10 x 1 film-coated tablets.

Lixiana 30 mg film-coated tablets are pink, round-shaped (8.5 mm diameter) and debossed with “DSC L30” on one side.
They come in blisters in cartons of 10, 14, 28, 30, 56, 60, 84, 90, 98 or 100 film-coated tablets or unit dose blisters in cartons of 10 x 1, 50 x 1, or 100 x 1 film-coated tablets.

Lixiana 60 mg film-coated tablets are yellow, round-shaped (10.5 mm diameter) and debossed with “DSC L60” on one side.
They come in blisters in cartons of 10, 14, 28, 30, 56, 60, 84, 90, 98 or 100 film-coated tablets or unit dose blisters in cartons of 10 x 1, 50 x 1, or 100 x 1 film-coated tablets.

Not all pack sizes may be marketed.

Marketing Authorisation Holder

Daiichi Sankyo Europe GmbH
Zielstattstrasse 48
81379 Munich
Germany

Manufacturer

Daiichi Sankyo Europe GmbH
Luitpoldstrasse 1
85276 Pfaffenhofen
Germany
For any information about this medicine, please contact the local representative of the Marketing Authorisation Holder:

**België/Belgique/Belgien**  
Daiichi Sankyo Belgium N.V.-S.A  
Tél/Tel: +32-(0) 10 48 95 95

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Daiichi Sankyo UK Ltd  
Tel: +44-(0) 1753 893 600

This leaflet was last revised in {MM/YYYY}.

Other sources of information
Detailed information on this product is available by scanning the QR Code below with a smartphone. The same information is also available on the following URL: www.dspatient.eu.

QR code to be included
Detailed information on this medicine is available on the European Medicines Agency web site: http://www.ema.europa.eu.
ANNEX IV

SCIENTIFIC CONCLUSIONS AND GROUNDS FOR THE VARIATION TO THE TERMS OF THE MARKETING AUTHORISATION(S)
Scientific conclusions

Taking into account the PRAC Assessment Report on the PSUR(s) for edoxaban, the scientific conclusions of the CHMP are as follows:

A number of cases of thrombocytopenia have been reported spontaneously and thrombocytopenia was uncommonly reported as an adverse event in pivotal clinical trials. Many of the spontaneously reported cases involved concomitant use of other medications and the causal role of edoxaban is difficult to establish, but equally in some cases the data are suggestive of a possible causal association. The MAH has already identified thrombocytopenia as a post-marketing adverse drug reaction (ADR) and the product information section 4.8 is being updated accordingly together with consequential changes to the Package Leaflet.

The CHMP agrees with the scientific conclusions made by the PRAC.

Grounds for the variation to the terms of the marketing authorisation(s)

On the basis of the scientific conclusions for edoxaban the CHMP is of the opinion that the benefit-risk balance of the medicinal product(s) containing edoxaban is unchanged subject to the proposed changes to the product information.

The CHMP recommends that the terms of the marketing authorisation(s) should be varied.