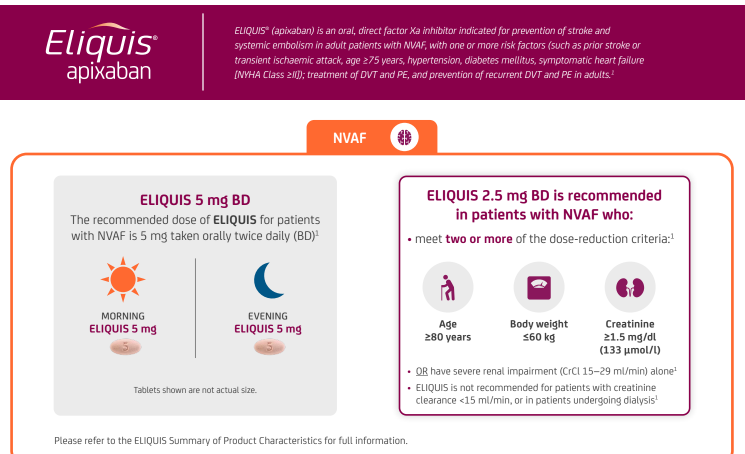
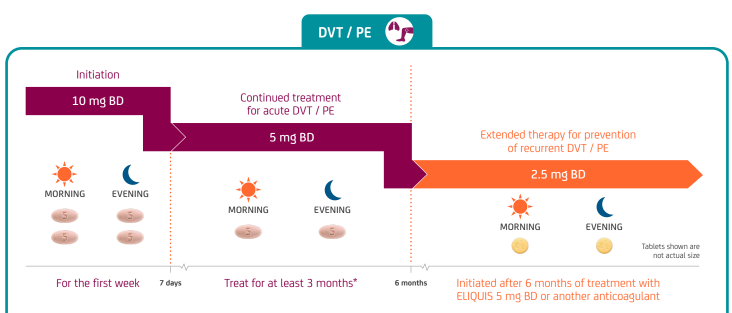
ELIQUIS® (apixaban) Dosing card

To assist you in selecting the right dose of ELIQUIS for the right patients. Refer to the Summary of Product Characteristics for full dosing considerations.





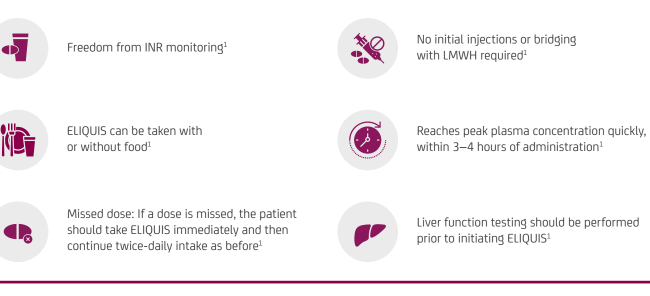
Duration of overall treatment should be individualised after careful assessment of the treatment benefit against the risk for bleeding.¹

⁶ As per available medical guidelines, short duration of treatment (at least 3 months) should be based on transient risk factors (e.g. recent surgery, trauma, immobilisation).¹



- No dose adjustments for the treatment of DVT / PE and prevention of recurrent DVT / PE (VTEt), based on age, weight or those with mild-to-moderate renal impairment¹
- ELIQUIS should be used with caution in patients with severe renal impairment (CrCl 15–29 ml/min) for the treatment of DVT / PE and prevention of recurrent DVT / PE.¹ ELIQUIS is not recommended in patients with CrCl <15 ml/min, or in patients undergoing dialysis¹

ELIQUIS IS A CONVENIENT, ORAL ANTICOAGULATION TREATMENT FOR PATIENTS



DOSING CONSIDERATIONS FOR PATIENTS WITH HEPATIC IMPAIRMENT

ELIQUIS is contraindicated in patients with hepatic disease associated with coagulopathy and clinically relevant bleeding risk.¹

ELIQUIS is not recommended in patients with severe hepatic impairment.¹

ELIQUIS should be used with caution in patients with mild or moderate hepatic impairment (Child Pugh A or B).¹ No dose adjustment is required in patients with mild or moderate hepatic impairment.¹

Patients with elevated liver enzymes alanine aminotransferase (ALT)/aspartate aminotransferase (AST) >2 x ULN or total bilirubin ≥1.5 x ULN were excluded in clinical trials. Therefore ELIQUIS should be used with caution in these patients.1

Prior to initiating ELIQUIS, liver function testing should be performed.¹

Please refer to the ELIQUIS Summary of Product Characteristics for full information.

BD = Twice Daily CrCl = Creatinine Clearance DVT = Deep Vein Thrombosis INR = International Normalized Ratio LMWH = Low Molecular Weight Heparin NVAF = Non-Valvular Atrial Fibrillation NYHA = New York Heart Association PE = Pulmonary Embolism TIA = Transient Ischaemic Attack ULN = Upper Limits of Normal

Reference

1. ELIQUIS® (apixaban) Summary of Product Characteristics. Available at www.medicines.org.uk.









ELIQUIS[®] (apixaban) PRESCRIBING INFORMATION United Kingdom

Consult Summary of Product Characteristics (SmPC) before prescribing

ELQUIS® (apixaban) PRESCRIBING INFORMATION UNITED TO PROVIDE THE ADVISOR OF THE The ormatics consider that the set of the se

be restarted as soon as possible. *Spinal/epidural anaesthesia or puncture*: Patients treated with antithrombotic agents for prevention of thromboembolic complications are at risk of developing an epidural or spinal haematoma which can result in long- term or permanent paralysis. The risk of these events may be increased by the post-operative use of indevelling epidural catheters or the concomitant use of medicinal products affecting haemostasis. Indevelling epidural catheters or the concomitant use of medicinal products affecting haemostasis. Indevelling epidural catheters or the concomitant use of medicinal products affecting haemostasis. Indevelling epidural catheters or the presence of the first dose of Eliquis. The risk may also be increased by traumatic or repeated epidural or spinal puncture. Patients are to be frequently monitored for signs and symptoms of neurological impairment (e.g., numbness or weakness of the legs, bowel or bladder dysfunction). If neurological compromise is noted, urgent diagnosis and treatment is necessary. Prior to neurovalial intervention the physician should consider the potential benefit intrathecial or epidural catheters. See SmPC for further details. *Haemodynamically unstable PE patients with active cancer* Patients with active cancer or Eliquis is not recommended as an alternative to unfractionated heparin in patients with pulmonary embolism who are haemodynamically unstable or may receive thrombolysis or pulmonazy embolectomy. Since the safety and efficacy of Eliquis when the one set abilished. *Patients with active cancer* can be at high risk of both venous thromboembolism and bleeding events. When aphaban is considered for DUT or PE treatment in cancer patients, dec., dec., and and administration section. *Itdextry patients with active cancer* and at a stry and Page. The eliquis is not commended with store inhibitors of the treatment of DUT and PE treatment to accempatient with active accompatient systemic treatment with storing inducers of both CYP3A4 and P-gg. Eliqui activated charcoal reduces Eliquis exposure. Also see contraindications and special warnings and precautions section; Consult SmPC (contraindications, special warnings and precautions section; Consult SmPC (contraindications, special warnings and precautions and drug pregancy, As a precautionary measure, it is preferable to avoid the use of apixaban during pregancy. As a precaution must be made whether to discontinue breast-feeding for the discontinue breast-feeding for the discontinue breast-feeding for the worman. UNDESIRABLE EFFECTS (SPC Section 4.8); increased risk of occult or overt bleeding from any tissue or organ, which may result in post haemorrhagi anaemia. The signs, symptoms, and severity will vary according to the location and degree or extent of the bleeding. Frequencies: common (≥ 1/100 to < 1/10); uncommon (≥ 1/1,000 to < 1/10,00); tev rare (< 1/10,000 to < 1/10,000; not known (cannot be estimated from the available data). Prevention of VTE in adult patients who have undergone elective hip or knee replacement surgery (VTEp): *Common*; anaemia; haemorrhage"; haematoma"; nausea; contusion. *Uncommon*; thrombocytopenia"; epistaxis"; haematochezia"; liver function test abnormal (including blood bilirubin increased'); haematuria"; specific haemorrhage such as gastrointestinal, abnormal waginal; urogenital", post procedural", would sceretion", incision site", operative". *Rare:* typersensitivity"; anaphylaxis"; haemotorysis"; gingival bleeding"; specific haemorrhage such as brain (encompassing intracranial, intraspinal), intra-abdominal", reprintage such as verit (including conjunctival), rectary such as respiratory tract, haemorrhage: nausea; gingival bleeding"; nausea; gingival bleeding"; anamatochezia"; hower for chaemorrhage such as verit (including conjunctival), gastrointestinal", rectary, haemorrhage'; haematorhage'; haematorhage';

Adverse events should be reported. Reporting forms and information can be found via: United Kingdom - The yellow card scheme at www.mhra.gov.uk/yellowcard, or search for MHRA Yellow Card in the Google Play or Apple App Store Adverse events should also be reported to Bristol-Myers Squibb via medical.information@bms.com or 0800 731 1736 (United Kingdom)

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📕 Uh Bristol Myers Squibb" 🛛 🔁 Pfizer

