Boehringer Ingelheim is pleased to provide you with this Educational Pack. which has been developed to give practical and relevant information on the appropriate use of Pradaxa<sup>®</sup>. The pack includes:

- Pradaxa<sup>®</sup> 150mg b.d Summary of Product Characteristics
- Pradaxa<sup>®</sup> 110mg b.d. Summary of Product Characteristics
- Presciber Guide this addresses recommendations for the use of Pradaxa<sup>®</sup> in order to minimize the risk of bleeding
- Patient Alert Card

To order additional copies of the Patient Alert Card please go to: www.pradaxa.co.uk/educationalpack

You can also order or download this Educational Pack.

## References:

Boehringer Ingelheim. Pradaxa® 150mg SPC. 2011.
Boehringer Ingelheim. Pradaxa® 110mg SPC. 2011.

## Prescribing Information (SPAF – UK) PRADAXA® (dabigatran etexilate)

Proscribing Information (SPAF – UK) PRADAXA\*\*\* (diabigatran etexilate) Capasies containing 110 mg or 150 mg obligatian etexilate (as mealate) Action: The measurement of dabigatran related anticcapulation mg be helpful to another measurement of dabigatran related anticcapulation mg be helpful to another measurement of dabigatran related anticcapulation mg be helpful to another measurement of dabigatran related anticcapulation mg be helpful to another measurement of dabigatran related anticcapulation mg be helpful to another measurement of dabigatran related anticcapulation mg be helpful to another measurement of dabigatran related anticcapulation mg be helpful to another measurement of dabigatran related anticcapulation mg be helpful to another measurement of dabigatran related anticcapulation mg be helpful to another measurement of dabigatran related anticcapulation mg be helpful to another measurement of dabigatran related anticcapulation mg be helpful to another measurement of dabigatran related anticcapulation mg be helpful to another measurement of dabigatran related anticcapulation mg be helpful to another measurement of dabigatran related anticcapulation mg be helpful to another measurement of dabigatran related anticcapulation mg be helpful to another measurement of dabigatran related anticcapulation mg be helpful to another measurement of dabigatran related anticcapulation mg be helpful to another measurement of dabigatran related anticcapulation mg be helpful to another mg bare and thereaft bare measurement of dabigatran related anticcapulation mg be helpful to another measurement of dabigatran related anticcapulation mg be helpful to another mg bare and thereaft bare measurement of dabigatran related anticcapulation mg be helpful to another mg bare and thereaft bare measurement of dabigatran related anticcapulated in the measurement of dabigatran related anticcapulated anticego supervises and the second of the physics in the date second to measurement of the helpful to another measurement of dabigatra 10 mg cansule twice daily: Pradaxa and veranamil should be taken at the same time. No dose adjustment required but close clinical surveillance in patients < 50 kg. Not recommended if liver enzymes > 2 Upper Limit of Normal (ULN). If switching from Accomparison of the parenteral anticoagulant wall 12 hours are no case about be given 0-2 switching from parenteral anticoagulants to Pradaxa then Pradaxa about be given 0-2 from the parenteral anticoagulant wall 12 hours are no case of continuous treatment, if switching from Pradaxa to WA administration together with Pradaxa than continuous treatment, if switching from Pradaxa to WA addist the stating from of the WA about about the stating of the treatment dd not appear to reduce the extent of administration together with Pradaxa than continuous treatment, if switching from WA are paratory and so that the stating of the treatment dd not appear to reduce the extent of administration together with Pradaxa than contained regrams of units whole with water, with or without food. Patients should be instructed not to open the whole with water, with or without food. Patients should be instructed not to open the administration together with area of SEC. Common (e. 1700, <170); an agencies and the patient indexisting and SEC. Common (e. 1700, <170); an agencies and the patient indexisting and SEC. Common (e. 1700, <170); an agencies and the patient indexisting and the patient indexisting in the attending and the state and SEC. Common (e. 1700, <170); an agencies and the patient indexisting and the patient indexisting and the state and SEC. Common (e. 1700, <170); an agencies and the state and SEC. Common (e. 1700, <170); an agencies and whole with antigether with stress and With and the state and SEC. Second and the statement and the statement app. (150 mg). Prescribers allowed and the statement app. (150 mg). Prescribers allowed and the statement app. (150 mg). An agencies and With and the statement app. (150 mg). An agencies and With and the statement app. (150 mg). An agencies and With and the statement app. (150 mg). An agencies and With and the statement app. (150 mg). An agencies and With and the statement app. (150 mg). An agencies and With and the statement app. (150 mg). An agenc whole wan water, while or window noot nearbits should be instructed into to get the capable as this may increase the risk of bledning, contraminations: hypersensitivity the teated for the prevention of stoke and SEE. Common (e 1/100, -1/10), anemina; to any component; severe renal impairment (O.C. < 30 m/min); active dinically epistaxis gastronitestinal haemorthage stabilitiations; dyspessia, nauses; significant bleeding; organic lesion at risk of bleeding; impairment of haemostasis; peritourological haemorthage (150 mg). Prescribers should consult the Summary of hepatic impairment or fiver disease expected to have any impact on survival. Phodul Characteristics for Unither information on side effects. **Pack stars and NNS** inepartie impairment of new basese expected to new any impact of survey, concomitant systemic ketoomarose, cyclosoprine, inconazole, laconitions. Warnings and Precautions: Not recommended if liver enzymes > 2 UUN Haemorrhagic risk. Close clinical surveillance (signs of bleeding or anaemia) is recommended throughout the treatment period, especially when haemorrhagic risk is increased or risk factors combined. Factors which may increase haemorrhagic risk: age ≥ 75 years; moderate renal impairment (CrCL 30 – 50 ml/min); P-glycoprotein inhibitor co-medication; body weight < 50 kg; acetylsalicylic acid (aspirin); NSAID; clopidogrel; diseases/procedures associated with a risk of bleeding such as coagulation disorders, thrombocytopenia or functional platelet defects, active ulcerative Gl disease, recent Gl bleeding, recen biopsy or majortrauma, recent ICH or brain, spinal or ophthalmic surgery, bacteria

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protease inhibitors including ritonavir and its combinations with other protease inhibitors avoid with P-gp inducers e.g. rifampicin, St John's wort, carbamazepine, phenytoir Dabigatran etexilate and dabigatran are not metabolised by cytochrome CYP450 sy Product unaradienties for further information on side effects, Pack steels and who price: 110mg 60 capsules 27:50 capsules 27:50 capsules 27:50 capsules 27:50 capsules 27:50 Manumbers: 110mg EUV/08/442/007 (Ro capsules) 150 mg EUV/08/442/011 (Ro capsules) Marketing Authorisation Holder: Boehringer Ingelheim International GmbH, D-55216 ingelheim am Rhein, Germany, Preschers should consult the Summary of Product Characteristics for full prescribing information. Prepared in July 2011

Adverse events should be reported. Reporting forms and information can be found at <u>www.yellowcard.gov.uk</u> Adverse events should also be reported to Boehringer Ingelheim Drug Safety on 0800 328 1627 (freephone).

Date of preparation: June 2011 Job code: DBG 2653

## **PRADAXA**<sup>®▼</sup> (DABIGATRAN ETEXILATE) EDUCATIONAL PACK

For prevention of stroke and systemic embolism in adult patients with nonvalvular atrial fibrillation with one or more of the following risk factors:

- Previous stroke, transient ischaemic attack or systemic embolism (SEE)
- Left ventricular ejection fraction <40%
- Symptomatic heart failure, ≥ New York Heart Association (NYHA) Class 2
- Age  $\geq$ 75 years
- Age ≥65 years associated with one of the following: diabetes mellitus, coronary artery disease or hypertension<sup>1,2</sup>