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®. The pack includes:

- Pradaxa® 150ma -Summary of Product Characteristics
- Pradaxa® 110mg -Summary of Product Characteristics
- Prescriber Guide this addresses recommendations for the use of Pradaxa® in order to minimise the risk of bleeding
- Patient Alert Card

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

You can also order or download this Educational Pack.

- Boehringer Ingelheim. Pradaxa* 150mg hard capsules Summary of Product Characteristics
 Boehringer Ingelheim. Pradaxa* 110mg hard capsules Summary of Product Characteristics

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

neoptasms at high risk of bleeding, recent brain or spinal injury, recent brain, spinal or optithalinic surper, recent intrazinal haemortange, known or suspected ecophiageal various, arterioverous mailformations, vasculair aneurysms or major intraspinal or intracerioral vascular abnormatiles, concomitant treatment with any other arterioragalants e.g. untractionated heparin (EHI), beyon interhorises fortispinarius etc), or all anticoagalants (warfain, intracolant, apixaban etc) except under the circumstances of switching therapy to or from Pradasa or when LPH is given at doses necessary to martinian an open central venous or arterial catheter, hepatic impairment or liver disease expected to have any impact on survivic, concomitant systemic ketocomizatio, spoisoprine, fizzonomici, barrollinis, directionated prospective fixer any size of the entire of the concentration of the complexity of the complexity of the complexity of the complexity of the Radional Confederations. Was included the control of the complexity of the Radional Confederations of the complexity of the Radional Confederation of the complexity of the Radional Confederation of the complexity of the Radional Confederation of the secondary of the recommendation of the teatment period, especially when haemortanging ricks labora confideration.

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



Date of preparation: February 2013 Job code: UK/DBG-121533(1)

PRADAXA® (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 ≥75 years
- Age ≥65 years associated with one of the following: diabetes mellitus, coronary artery disease or hypertension^{1,2}