NATIONAL INSTITUTE FOR HEALTH AND CLINICAL EXCELLENCE

Single Technology Appraisal (STA)

Dabigatran for the prevention of deep vein thrombosis after hip or knee replacement surgery in adults

Matrix of consultees and commentators

Consultees	Commentators (no right to submit or appeal)
Manufacturers/sponsors	General
Boehringer-Ingelheim (dabigatran)	 Department of Health, Social Services and Public Safety for Northern Ireland
Patient/carer groups	NHS Quality Improvement Scotland
Anticoagulation Europe	Wife Quality improvement occurance
DVT Awareness Campaign	Possible comparator manufacturer(s)
Lifeblood: The Thrombosis Charity	GlaxoSmithKline (fondaparinux sodium)
Professional groups	 Pfizer Ltd (dalteparin sodium)
 British Association for Surgery of the Knee 	Sanofi-Aventis Ltd (enoxaparin sodium)
 British Haematology Society 	Relevant research groups
 British Hip Society 	■ (None)
 British Orthopaedic Association 	
British Thoracic Society	Evidence Review Group
 British Society for Haemostasis and Thrombosis 	 School of Health & Related Research Sheffield (ScHARR)
Royal College of Nursing	 National Coordinating Centre for Health
Royal College of PathologistsRoyal College of Physicians	Technology Assessment
Troyal College of Fifty Stolaris	Associated Guideline Groups
<u>Others</u>	National Collaborating Centre for Acute
Department of Health	Care
 Welsh Assembly Government 	
	Associated Public Health Groups (None)

NICE is committed to promoting equality and eliminating unlawful discrimination.

Please let us know if we have missed any important organisations from the lists contained within the matrix and which organisations we should include who have a particular focus on relevant equality issues.

PTO FOR DEFINITIONS OF CONSULTEES AND COMMENTATORS

National Institute for Health and Clinical Excellence Matrix of consultees and commentators for the appraisal of dabigatran for the prevention of deep vein thrombosis after hip or knee replacement surgery in adults Issue date: January 2008

Definitions:

Consultees

Organisations that accept an invitation to participate in the appraisal; the manufacturer(s) or sponsor(s) of the technology; national professional organisations; national patient organisations; the Department of Health and the Welsh Assembly Government and relevant NHS organisations in England.

The manufacturer/sponsor of the technology is invited to make an evidence submission, respond to consultations and has the right to appeal against the Final Appraisal Determination (FAD).

All non-manufacturer/sponsor consultees are invited to submit a statement¹, respond to consultations, nominate clinical specialists or patient experts and have the right to appeal against the Final Appraisal Determination (FAD).

Commentators

Organisations that engage in the appraisal process but that are not asked to prepare an evidence submission or statement, are able to respond to consultations and they receive the FAD for information only, without right of appeal. These organisations are: manufacturers of comparator technologies; NHS Quality Improvement Scotland; the relevant National Collaborating Centre (a group commissioned by the Institute to develop clinical guidelines); other related research groups where appropriate (for example, the Medical Research Council [MRC], National Cancer Research Institute); other groups (for example, the NHS Confederation, NHS Information Authority and NHS Purchasing and Supplies Agency, and the *British National Formulary*.

All non-manufacturers/sponsors commentators are invited to nominate clinical specialists or patient experts.

Evidence Review Group (ERG)

An independent academic group commissioned by the NHS Research and Development Health Technology Assessment Programme (HTA Programme) to assist the Appraisal Committee in reviewing the manufacturer/sponsor evidence submission to the Institute.

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¹ Non manufacturer consultees are invited to submit statements relevant to the group they are representing.