GUIDELINES FOR FOLLOW UP OF IMPLANTABLE CARDiac DEVICES FOR CARDiac RHhythm MANAGEMENT
– October 2008

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INTRODUCTION

Guidelines for pacemaker follow up were first published in Heart in November 1996 as the result of a British Pacing and Electrophysiology Group (BPEG) policy conference. These were agreed by consensus and have been the primary source of guidance for the management of implanted devices within the UK to date.

The continuing evolution of device technology has resulted in the production of a range of devices capable of treating bradycardias, complex cardiac tachyarrhythmias, and heart failure management. These devices have multiple modalities and programmable features. The challenge of these treatments lies not only in the implantation of the devices but also in comprehensive follow-up of the implanted devices as part of the lifelong management of the patient. As the number and variety of implanted devices increase so does the burden of follow-up and the knowledge required to optimise and troubleshoot their use. This is compounded by the increasing volume of data provided by devices, and the increasing sophistication of programming therapy and detection algorithms. As a result of this increasing complexity, inappropriate or incorrect use of these algorithms or other errors with aspects of device programming may result in serious harm to the patient.

Device follow-up clinics now encompass various types of devices ranging from single/dual chamber bradycardia devices, atrial tachycardia devices and implantable cardioverter defibrillators, to multi-chamber cardiac resynchronisation devices incorporating new modalities such as impedance monitors for assessing heart failure. Although the original guidelines stated that pacemaker clinics should be the responsibility of a lead physician, the practice of device management follow-up in the UK is now almost solely led and practiced by Cardiac Physiologists (CPs), and in some cases by specialist nurses. It is vital therefore that clinical governance mechanisms and lines of clinical responsibility are clearly established for all follow-up clinics.
Device follow-up remains the clinical responsibility of the Consultant Cardiologist (consultant physician with specialist interest) in charge of the device follow-up service – although it is a Cardiac Physiologist run service. Physicians providing such a service should have the required knowledge to do so. Ideally, such physicians would be current holders of (or working towards) a recognised pacing qualification such as certificate of accreditation with Heart Rhythm UK (HRUK, previously BPEG exam), European Heart Rhythm Association (EHRA) or International Boards of Heart Rhythm Examiners (IBHRE, formerly North American Society of Pacing and Electrophysiology (NASPE)).

The following guidelines for Cardiac Physiologist led clinics have been drawn up by HRUK and the Society for Cardiological Science and Technology (SCST), both of which are affiliated groups of the British Cardiovascular Society.

These groups have been instrumental in guiding device follow up training since the previous policy conference conducted by the British Pacing and Electrophysiology Group (now HRUK).

**Section A Bradycardia Pacemaker Follow up Clinics**

There should be a clearly defined protocol documenting the lines of communication and support between the lead Cardiac Physiologist for the bradycardia pacemaker follow-up service and the Consultant Cardiologist (consultant physician with specialist interest) responsible for the onsite service to ensure that clinical governance requirements are met. The lead Cardiac Physiologist for bradycardia pacemaker follow-up services at non-implanting hospitals must also have strong links with the lead Cardiac Physiologist and Consultant Cardiologist at the implant centre.

The lines of clinical responsibility must be clearly defined in the local trust policy. Trusts delivering bradycardia pacemaker follow-up services have a responsibility to ensure appropriate arrangements are in place to cover clinic activity (elective or urgent).

**Bradycardia Pacemaker Follow-up Clinic Objectives:**

1. To optimise the pacing system to the individual patient needs whilst maximising generator life. Safety must be paramount whilst manufacturer guidance and HRUK recommendations should also be taken into account.
2. To identify any abnormalities in the pacemaker system and complications of the therapy in order to ensure prompt treatment.
3. To assess battery status to predict end-of-life (EOL) of the pulse generator in order to permit timely elective generator replacement.
4. To provide patient and family support and education.
5. To ensure that the patient’s experience is as safe, comfortable and reassuring as possible.
6. To ensure that safe and accurate measurements are made of device and lead function and that accurate records of each visit are kept. Staff leading the clinic must be able to recognise problems and complications and make the appropriate changes or recommendations.
7. To monitor the device implant site and manage any risk of infection.
8. To maximise clinical safety and efficiency in line with clinical governance requirements.
9. To regularly review patients in line with local, manufacturer and National guidelines.
10. To implement relevant advisories from device manufacturers and the Medicines and Healthcare products Regulatory Agency (MHRA) guidelines and advice.
11. To notify the MHRA and manufacturer of any problems arising with devices or leads.
12. To be aware of the need to identify clinical problems and refer patients for immediate or deferred medical care appropriately in line with local policy.
13. To provide accurate and complete communication about patient-device interaction and appropriate functionality to GPs and other relevant health professionals.

Suggested Procedure for Referrals to a Bradycardia Pacemaker Service Follow-Up Clinic at non-implant centres

On receipt of referral from implant physician/centre the patient is registered and scheduled for review. The referral information must include:

- Patient name and address and telephone number
- Patient GP details
- Hospital / H&C number
- Date of birth
- Referring consultant
- Pacemaker type and parameters
- Pacemaker lead models and serial numbers
- Most recent threshold, battery voltage and lead impedance evaluation results
- Patient mobility
- Cross infection issues
- Indications for implant
- Medication details.

Suggested appointments schedule

- Yearly for pacemakers implanted for less than 7-10 years depending on the manufacturers recommendation and expected battery longevity.
- 6 monthly for implants exceeding 7-10 years until ERI is reached.
- 3-6 monthly for Devices that exceed the manufacturers suggested longevity or show decline in battery life.
- At Cardiac Physiologists discretion for devices that require closer monitoring e.g. programming/lead issues.

Reports

- A report is generated by Cardiac Physiologists in charge of the follow-up clinic and all parameters and clinical details are documented in department's database and/or the patient's pacemaker notes.
- A copy of the report or a letter is sent to the patient’s general practitioner and the referring hospital where appropriate.
The implant centre need only be contacted when seeking additional advice or when making a referral to the implanting physician.

**Staffing: Qualifications and Training**

Ideally the clinic should be manned with two staff, one of who meets the lead role competencies. The Second staff member can be undergoing training.

**Lead Cardiac Physiologist**

- A qualified Cardiac Physiologist (BSc Clinical Physiology or equivalent)
- Evidence of post-graduate training in cardiac rhythm management techniques, e.g. holds appropriate Certification of Accreditation HRUK or IBHRE
- Hold current ILS or ALS accreditation
- Evidence of Continual Professional Development (CPD) in cardiac rhythm management
- Perform minimum of 100 bradycardia pacemaker system follow-up review procedures per year
- Attend local implant centre regularly and not less than twice per annum to remain familiar with evolving technology (CP’s leading Follow-Up Clinic at non implant hospitals)
- Demonstrate high level of understanding and knowledge of the full range of diagnostic cardiac investigations

**Cardiac Physiologist**

- A qualified Cardiac Physiologist (BSc Clinical Physiology or equivalent)
- Holds a current ILS accreditation
- Has a proven understanding of bradycardia pacemaker implant procedures
- Has a proven knowledge of bradycardia pacemaker technology
- Has current CPD by attending relevant recognised training study days

**Equipment and Other Essential Requirements**

A wide range of equipment is essential within the clinic or immediate vicinity of the clinic with access to further cardiac investigations (which need not necessarily be on site). These are listed below.

Equipment essential in the Pacemaker clinic (or in the immediate vicinity):
- 12-Lead electrocardiograph (ECG) machine with real time recording
- An appropriate range of manufacturer programmers (with appropriate documentation for use of each specific model)
- Emergency ‘crash’ trolley and defibrillator with integrated pacing function.
- Magnet
- Wound treatment pack
- Telephone and/or arrest call button
- Callipers, rate ruler etc
- Data management system/patient notes
- Sharps box
• Oxygen, suction and relevant adjuncts.

There should also be access to MHRA Pacemaker adverse incident reporting forms.

Investigations to which the cardiac physiologist should have referral access:
• X-Ray facilities
• Ambulatory ECG Recording
• Echocardiogram.

Cardiac investigations to which it may be desirable to have referral access:
• Exercise Stress Testing
• Head-up Tilt-Table Testing.

**Access for rapid referral of any patient needing urgent admission should also be available.**

**Section B ICD/CRT Device Follow up Clinics**

There should be a clearly defined protocol documenting the lines of communication and support between the lead Cardiac Physiologist for the implantable cardio defibrillator (ICD) and cardiac resynchronisation therapy (CRT) follow-up service and the Consultant Cardiologist responsible for the onsite service to ensure that clinical governance requirements are met. ICD and CRT follow-up clinics should not be undertaken without a designated physician available onsite.

The lines of clinical responsibility must be clearly defined in the local trust policy. Trusts delivering ICD/CRT device follow-up services have a responsibility to ensure appropriate arrangements are in place to cover clinic activity (elective or urgent).

**ICD/CRT Device Clinic Follow Up Aims and Objectives**

1. To optimise the system to provide delivery of optimal therapy for the individual patient needs whilst maximising generator life. Safety must be paramount whilst manufacturer guidance and HRUK recommendations should also be taken into account.
2. To identify any abnormalities in the ICD/CRT system and any complications of the therapy in order to ensure prompt treatment.
3. To assess battery status to predict end-of-life (EOL) of the pulse generator in order to permit timely elective generator replacement.
4. To provide patient and family support and education together with any other healthcare professionals involved in the patient’s management.
5. To ensure that the patient’s experience is as safe, comfortable and reassuring as possible.
6. To ensure that safe and accurate measurements are made and that accurate records of each visit are kept. Staff leading the clinic must be able to identify problems and complications and make the appropriate changes or recommendations.
7. To monitor the device implant site and manage any risk of infection.
8. To maximise clinical safety and efficiency in line with clinical governance requirements.
9. To regularly review patients in line with local, manufacturer and National guidelines.
10. To implement relevant advisories from device manufacturers and the Medicines and Healthcare products Regulatory Agency (MHRA) guidelines and advice.
11. To notify the MHRA and manufacturer of any problems arising with devices or leads.
12. To be aware of the need to identify clinical problems and refer patients for immediate or deferred medical care appropriately in line with local policy.
13. To provide accurate and complete communication about patient-device interaction and appropriate functionality to GPs and other relevant health professionals.
14. To monitor any counters of intra-cardiac events or episodes to ensure that appropriate therapy is being delivered and to minimise patients symptoms where necessary.

Staffing: Qualifications and Training

Ideally the clinic should be manned with two staff, one of who meets the lead role competencies. The Second staff member can be undergoing training.

Lead Cardiac Physiologist

- A qualified Cardiac Physiologist (BSc Clinical Physiology or equivalent)
- Evidence of post-graduate training in cardiac rhythm management techniques e.g. holds appropriate Certification of Accreditation HRUK or IBHRE
- Hold current ILS or ALS accreditation
- Evidence of Continual Professional Development (CPD) in cardiac rhythm management
- Perform minimum of 100 ICD/CRT device follow-up review procedures per year
- Demonstrate high level of understanding and knowledge of the full range of diagnostic cardiac investigations
- Must attend an ICD/CRT device course, which has been accredited by HRUK or SCST at least once per year

In some cases it may be appropriate that with the relevant training and competencies that the lead Cardiac Physiologist may take responsibility for titrating drugs for heart failure optimisation.

Cardiac Physiologist

- A qualified Cardiac Physiologist (BSc Clinical Physiology or equivalent)
- Holds current ILS accreditation
- Proven understanding and experience in bradycardia pacemaker and ICD/CRT device implant procedures
- Has a proven knowledge of bradycardia pacemaker and ICD/CRT device technology
- Has a proven experience in bradycardia pacemaker follow up clinics
- Has current CPD by attending relevant recognised training study days
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Section C Transmitted/Remote Device follow-up

With a rapidly evolving technology, telephone transmission is likely to change the face of device follow up and management in the very near future. There are already several methods of telephonic transmission available and more are currently in development.

Each company may use different technology and this will add further complexity to device follow-up services. However it is anticipated that there will be added benefits in reduced patient journeys and unnecessary hospital admissions and therefore allowing more time available for complex device follow-up by experienced Cardiac Physiologists.

The Cardiac Physiologist supervising such a remote service will therefore require the same level of expertise and training of the Cardiac Physiologist leading a clinic attended directly by patients.
Guidance will be taken from manufacturers and the European taskforce group on regulations relating to Remote device follow up and Trust protocols, agreed by the responsible Consultant Cardiologist/Lead Cardiac Physiologist must be in place.

Clinic Procedures

All device follow up clinics should work to a standard procedure/protocol. This may be locally developed but should incorporate the minimum requirements set out in these guidelines.

A procedure for device follow-up should include the following where possible:

- Recording of an ECG rhythm strip to verify device function and monitoring this throughout the check
- Maintenance of device function throughout check
- Identification of the device and leads from the patient records
- Initial interrogation of the device and recording of any relevant information
- Assessment of device battery status and comparison with previous records
- Safe testing of device and lead status including thresholds for sensing and capture as well as impedance measurements
- Assessment of diagnostics, events and appropriate counters/histograms for rate assessment and appropriate function
- Appropriate troubleshooting for complications/problems using other investigations where necessary
- Appropriate reprogramming of the device to ensure that optimal settings for clinical outcomes are provided for the patient
- Recording all the above
- Checking final settings and ensuring that any changes have been fully and appropriately documented and checked to ensure patient safety
- Appropriate scheduling of the next appointment or referral
- Ensuring that device registration has been undertaken and that all patients have their registration/ID cards and all appropriate information.

Special thanks to the following individuals: Ian Culshaw, Chris Eggett, Jacqui Howard, Tony McEntee and Geraldine McParland for their contribution towards the production of this document.


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