How are arrhythmias detected by implanted cardiac devices managed in Europe? Results of the European Heart Rhythm Association Survey

Derick Todd¹*, Antonio Hernandez-Madrid², Alessandro Proclemer³, Maria Grazia Bongiorni⁴, Heidi Estner⁵, and Carina Blomström-Lundqvist⁶, Scientific Initiative Committee, European Heart Rhythm Association

¹Institute of Cardiovascular Medicine and Science, Liverpool Heart and Chest Hospital, Thomas Drive, Liverpool L14 3PE, UK; ²Cardiology Department, Ramon y Cajal Hospital, Alcalá University, Carretera Colmenar Viejo, Madrid, Spain; ³Division of Cardiology, University Hospital S. Maria della Misericordia, IRCAB Foundation Udine, Udine, Italy; ⁴2nd Cardiology Department, University Hospital of Pisa, Pisa, Italy; ⁵Medizinische Klinik, Ludwig-Maximilians-Universitat, Campus Großhadern, Munchen, Germany; and ⁶Department of Cardiology, Institution of Medical Science, Uppsala University, Uppsala 75185, Sweden

Received 29 July 2015; accepted after revision 19 August 2015

The management of arrhythmias detected by implantable cardiac devices can be challenging. There are no formal international guidelines to inform decision-making. The purpose of this European Heart Rhythm Association (EHRA) survey was to assess the management of various clinical scenarios among members of the EHRA electrophysiology research network. There were 49 responses to the questionnaire. The survey responses were mainly (81%) from medium–high volume device implanting centres, performing more than 200 total device implants per year. Clinical scenarios were described focusing on four key areas: the implantation of pacemakers for bradyarrhythmia detected on an implantable loop recorder (ILR), the management of patients with ventricular arrhythmia detected by an ILR or pacemaker, the management of atrial fibrillation in patients with pacemakers and cardiac resynchronization therapy devices and the management of ventricular tachycardia in patients with implantable cardioverter-defibrillators.

Keywords
Implantable loop recorder • Pacemaker • Implantable cardiac-defibrillator • EHRA survey • EP wire

Introduction

Over 250 000 devices are implanted annually in Europe,¹ and all of these devices have the ability to store details of arrhythmias, either as patient-activated events or as automatic recordings within pre-programmed criteria (normally heart rate). In many patients, additional arrhythmias to the original device indication will be detected and require management. There are no international guidelines on how to manage such arrhythmias. The purpose of this European Heart Rhythm Association (EHRA) survey was to assess how physicians within the EHRA electrophysiology (EP) research network managed a series of clinical scenarios often seen within device follow-up.

Methods

Participating centres

This survey is based on a questionnaire sent via the internet to the EHRA EP research network centres. Overall, 46 institutions responded, with 2 replies from 3 centres (all in the UK). There was a wide geographical distribution from 17 countries (19 centres in the UK, 5 in Spain, 3 in Italy and Denmark, 2 in Armenia, Austria, and Georgia, and 1 centre in each from Belgium, Brazil, Estonia, Greece, Luxembourg, The Netherlands, Norway, Portugal, Romania, and Serbia). The majority of centres (78%) were university hospitals, with 3 private hospitals (6%) and 8 (16%) other types of hospital. Hospital procedure numbers are shown in Table 1. Most centres (81%) were high volume implanting (>200 per annum) centres.

Bradyarrhythmia detected by implantable loop recorders

The first scenario was that of a patient with a history of syncope, a normal 12-lead electrocardiogram (ECG) and a structurally normal heart on echocardiography. The issue we explored was the threshold for pacemaker implantation based on the duration of monitored asystole and the presence or absence of symptoms. It was clear that symptoms were important in the decision to implant a pacemaker. In the absence of symptoms, only 7% of respondents would implant a pacemaker for a
3 s pause, whereas 72% would implant a pacemaker for a symptomatic 3 s pause, and a further 18% for a 5 s pause. In the absence of symptoms a pause of duration of 5 s was considered an indication for pacemaker by 40% of centres, whereas 23% would require a pause to be >10 s and 30% indicated that they would not implant a pacemaker for any asymptomatic pause.

### Table 1 Device implant numbers and types per centre

<table>
<thead>
<tr>
<th>Implant numbers</th>
<th>0</th>
<th>1–49</th>
<th>50–99</th>
<th>100–199</th>
<th>&gt;200</th>
</tr>
</thead>
<tbody>
<tr>
<td>All devices</td>
<td>0</td>
<td>0</td>
<td>3</td>
<td>3</td>
<td>40</td>
</tr>
<tr>
<td>Pacemakers</td>
<td>0</td>
<td>1</td>
<td>5</td>
<td>9</td>
<td>31</td>
</tr>
<tr>
<td>Loop recorders</td>
<td>7</td>
<td>21</td>
<td>11</td>
<td>7</td>
<td>0</td>
</tr>
<tr>
<td>ICD</td>
<td>1</td>
<td>6</td>
<td>12</td>
<td>12</td>
<td>15</td>
</tr>
<tr>
<td>CRT</td>
<td>2</td>
<td>14</td>
<td>15</td>
<td>9</td>
<td>6</td>
</tr>
</tbody>
</table>

CRT, cardiac resynchronisation therapy; ICD, implantable cardioverter-defibrillator.

### Broad complex tachycardia detected by implantable loop recorders

For implantable loop recorder (ILR) detected broad QRS complex tachycardia (BCT), we explored two scenarios for a patient with a history of syncope. In the first scenario, the patient had a normal 12-lead ECG and a structurally normal heart on echocardiography. For ‘normal heart’ ventricular tachycardia (VT), there was virtual universal agreement (93% of respondents) that an implantable cardioverter-defibrillator (ICD) was not indicated for an asymptomatic run of non-sustained VT at 200 bpm. However, if the patient had been symptomatic, 35% of centres indicated that they would consider an ICD implant, although 65% of centres still did not consider an ICD in such a case. Of the 35% who would consider an ICD for a patient with a normal heart and history of syncope, 12% would consider it for a 3 s run of BCT, 9% for a 5 s run, and 14% for a run >10 s.

In the second scenario, the patient with a history of syncope had structural heart disease, defined as a previous anterior myocardial infarction and an ejection fraction (EF) of 40% on echocardiography (chosen so as not to overlap with current primary prevention ICD indications). Slightly more than half of the centres (53%) did not feel an ICD was indicated for asymptomatic BCT, although a significant proportion (28%) felt that a 3 s run of BCT was enough to proceed to ICD implantation. For symptomatic BCT, 65% of respondents would implant an ICD, although there was also a wide variation in practice with 40% feeling an ICD was indicated for a 3 s episode, 12% for a 5 s episode, and 14% for a 10 s episode. However, 35% of centres still did not feel an ICD was indicated even for an episode of BCT >10 s with symptoms, even with an EF of 40%.

For pacemaker-detected tachycardia, we explored six scenarios, all in patients with underlying complete heart block, to strongly imply that any ventricular high-rate episodes were VT. In the first two scenarios, the patient was asymptomatic and had either an EF of 50 or 40% on echocardiography. We specified a rate of 185 bpm lasting 10 s. When EF was 50% and the patient was asymptomatic, no respondents chose to upgrade to an ICD, although most (74%) would start a β-blocker and investigate the patient’s coronary artery status (58%). A significant proportion of centres (19%) take no action in this situation. In the scenario where the patient had symptoms and an EF of 40%, upgrade to ICD was considered by 28% of centres, with similar proportions starting β-blocker (77%), and investigating coronary status (65%). Only 5% of centres took no action.

In the next four scenarios, we defined the high-rate episode as 200 bpm lasting 15 s. We explored how left ventricular function and symptoms affected management. In the first comparison, the patient had normal left ventricular function and we assessed how management was affected by symptoms. In the absence of symptoms, physicians were considerably more conservative, mostly advising β-blocker (71%) and coronary investigation (61%) but rarely an ICD (7%). If symptomatic, a β-blocker was recommended by 93% and coronary investigation by 71%. The presence of symptoms also served as a trigger for recommending an ICD for 20% (symptoms were not defined). Ablation was recommended by 15% of respondents.

When the same questions were asked but the scenario changed to a patient with an EF of 40%, the use of ICD increased significantly to 34% for asymptomatic episodes, and 46% in the presence of symptoms. For both groups the vast majority advised a β-blocker and coronary investigation. Ablation was still a favoured option in the presence of symptoms for 12% of centres.

### Atrial fibrillation detected by implanted devices

We explored the two key elements of atrial fibrillation (AF) management. First we explored the threshold for anticoagulation dependent on the duration of device-detected atrial arrhythmia and the patient’s CHA2DS2-VASc score. Second, we asked about rate vs. rhythm control management strategies in patients with cardiac resynchronisation therapy defibrillators (CRT-Ds).

The scenarios we explored involved patients with dual-chamber pacemakers, implanted for sinus node disease, coming for a routine 6-month follow-up appointment. We wanted to understand how respondents’ threshold for recommending anticoagulation would vary according to the CHA2DS2-VASc score, and the number and duration of device-detected AF events. The results are shown in Figure 1 and indicate that the threshold for initiation of anticoagulation is highly variable, but in general that the higher the CHA2DS2-VASc score, and the longer and more frequent AF episodes are, the more likely cardiologists are to recommend anticoagulation.

We also wanted to assess strategies for rate control vs. rhythm management in patients with CRT devices with both symptomatic and asymptomatic device-detected AF. The results are shown in Figure 2. There was a clear trend towards lower use of amiodarone in younger patients, and to a higher use of ablation to control AF in patients who are younger and symptomatic.

### Management of a first ICD shock for monomorphic ventricular tachycardia

In the final clinical scenario, we asked how respondents would manage a patient with a history of ischaemic heart disease, and a primary prevention ICD, who presented following a first ICD shock for monomorphic ventricular tachycardia (VT), there was virtual universal agreement that an implantable cardioverter-defibrillator (ICD) was indicated for asymptomatic BCT, although a significant proportion (28%) felt that a 3 s run of BCT was enough to proceed to ICD implantation. For symptomatic BCT, 65% of respondents would implant an ICD, although there was also a wide variation in practice with 40% feeling an ICD was indicated for a 3 s episode, 12% for a 5 s episode, and 14% for a 10 s episode. However, 35% of centres still did not feel an ICD was indicated even for an episode of BCT >10 s with symptoms, even with an EF of 40%.

For pacemaker-detected tachycardia, we explored six scenarios, all in patients with underlying complete heart block, to strongly imply that any ventricular high-rate episodes were VT. In the first two scenarios, the patient was asymptomatic and had either an EF of 50 or 40% on echocardiography. We specified a rate of 185 bpm lasting 10 s. When EF was 50% and the patient was asymptomatic, no respondents chose to upgrade to an ICD, although most (74%) would start a β-blocker and investigate the patient’s coronary artery status (58%). A significant proportion of centres (19%) take no action in this situation. In the scenario where the patient had symptoms and an EF of 40%, upgrade to ICD was considered by 28% of centres, with similar proportions starting β-blocker (77%), and investigating coronary status (65%). Only 5% of centres took no action.

In the next four scenarios, we defined the high-rate episode as 200 bpm lasting 15 s. We explored how left ventricular function and symptoms affected management. In the first comparison, the patient had normal left ventricular function and we assessed how management was affected by symptoms. In the absence of symptoms, physicians were considerably more conservative, mostly advising β-blocker (71%) and coronary investigation (61%) but rarely an ICD (7%). If symptomatic, a β-blocker was recommended by 93% and coronary investigation by 71%. The presence of symptoms also served as a trigger for recommending an ICD for 20% (symptoms were not defined). Ablation was recommended by 15% of respondents.

When the same questions were asked but the scenario changed to a patient with an EF of 40%, the use of ICD increased significantly to 34% for asymptomatic episodes, and 46% in the presence of symptoms. For both groups the vast majority advised a β-blocker and coronary investigation. Ablation was still a favoured option in the presence of symptoms for 12% of centres.

### Discussion

The results of this survey indicate that there is a significant variation in the management of device-detected arrhythmias. It was not...
possible to assess how this varied geographically due to the large number of countries with only a single responding centre.

Recommendations for pacemaker implantation are highly dependent on the underlying rhythm disturbance. It was not possible from our questions to understand what form of bradycardia respondents were considering in their answers. Asymptomatic pauses of 3 s detected by an ILR would certainly be a pacemaker indication in a patient with a prior history of syncope if there was documentation of atrioventricular block, or the patient had underlying sinus node disease. However, if the ILR was implanted for a suspicion of vasovagal syncope, the results of the ISSUE-3 (International Study on Syncope of Uncertain Etiology) study would suggest that pacing should be reserved for asymptomatic pauses $>6$ s, and symptomatic pauses $>3$ s. It is interesting that 30% of centres would not

---

**Figure 1** Physician recommendation of anticoagulation for four scenarios. (A) CHA2DS2-VASc score of 2–3, single atrial high-rate event. (B) CHA2DS2-VASc score of 2–3, multiple ($>2$) AHREs. (C) CHA2DS2-VASc score of 4, single atrial high-rate event. (D) CHA2DS2-VASc score of 4, multiple ($>2$) atrial high-rate events.

**Figure 2** Atrial fibrillation management in CRT-D patients (single episode of 72 h). (A) The 78-year-old patient with an asymptomatic 72 h episode. (B) The 78-year-old patient with a symptomatic 72 h episode. (C) The 53-year-old patient with an asymptomatic 72 h episode. (D) The 53-year-old patient with a symptomatic 72 h episode.
The increased risk of stroke, at an approximate relative risk of \(2.3,4,10\) is not as high as the five times higher risk noted for persistent AF.\(^6\) The variation in practice is therefore quite understandable as at this time there are also no published randomized trials confirming a benefit for anticoagulation for device-only detected AF, although trials are underway (e.g. ARTESiA, \(https://clinicaltrials.gov/ct2/show/NCT01938248\)).

Perhaps, surprisingly our results show that a rhythm control policy either with drugs or with ablation was favoured in the majority of patients with 80% of respondents recommending this (either in the form of amiodarone, switching to sotalol, or using AF ablation) for the 78-year-old patient and 87.5% for the 53-year-old patient with a CRT-D. It was interesting to understand that despite the data from AFFIRM (Atrial Fibrillation Follow-Up Investigation of Rhythm Management)\(^12\) of equivalence in outcomes between rate and rhythm control strategies in AF, 80% of respondents still prefer rhythm control in patients with a CRT-D device, even in the absence of symptoms. It is likely that this reflects a belief that such patients are more prone to deterioration in heart failure with the onset of AF. We may have expected that atrioventricular node ablation would be used more widely, than the observed 20%, in these patients.

Finally, the management of non-sustained VT resulting in an ICD shock is influenced by the patients age with amiodarone preferred in older patients.

**Limitations**

By their very nature short clinical scenarios cannot cover every aspect of the patient’s history and clinical findings. In addition, not all possible management plans can be provided as an option. The main limitations in the scenarios were that we did not describe symptoms and therefore these were open to interpretation. In addition, we did not offer the option of an electrophysiological study to assess inducibility of VT as an option for patients with an EF of 40%.

**Acknowledgements**


**Conflict of interest:** none declared.

**References**

Pacemaker-mediated tachycardia in an unconventional resynchronization device

Vedran Velagić*, Richard Matasić, and Maja Čikes

Department of Cardiovascular Medicine, University of Zagreb School of Medicine and University Hospital Centre Zagreb, Kiliptićeva 12, Zagreb 10000, Croatia

* Corresponding author. Tel: +385 91 7929284; fax: +385 1 2367512, E-mail address: vvelagic@gmail.com

A 67-year-old male with dilated cardiomyopathy and permanent atrial fibrillation was admitted due to decompensated heart failure. Standard medical therapy failed to control high ventricular rates which aggravated the heart failure. Therefore, atrio-ventricular-node ablation was performed and a biventricular pacemaker was implanted. Instead of a CRT device, we implanted a DDD pacemaker with the left ventricular (LV) lead in the atrial channel. After the implant, frequent paroxysms of wide QRS tachycardia with the rate of 130 b.p.m. occurred (Figure A).

Interrogation of the device revealed that the tachycardia cycle length was exactly at the upper tracking rate limit, and the intermittent sensing of T waves in the atrial channel was detected (Figure B). When a sensed event occurs after the post-ventricular atrial refractory period (PVARP), right ventricle pacing is triggered. That explains the different morphology of the tachycardia QRS complex. High standard sensing parameters in the atrial channel, combined with high amplitude T waves detected by the LV lead, caused an endless-loop tachycardia. The problem was solved by decreasing the sensitivity in the atrial channel and by prolonging the PVARP.

This off-label application of DDD devices can result with an unusual mechanism of a pacemaker-mediated tachycardia. DDIR mode can also be used to avoid this problem.

The full-length version of this report can be viewed at: http://www.escardio.org/Guidelines-&-Education/E%2Elearning/Clinical-cases/Electrophysiology/EP-Case-Reports.