Ventricular pacemaker upgrade: experience, complications and recommendations

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Abstract
Objective—To assess outcomes of pacemaker upgrade from single chamber ventricular to dual chamber.
Design—Retrospective analysis of patients undergoing the procedure.
Setting—Specialist cardiothoracic unit.
Patients—44 patients (15 female, 29 male), mean (SD) age at upgrade 68.2 (12.9) years.
Interventions—Upgrade of single chamber ventricular to dual chamber pacemaker.
Main outcome measures—Procedure duration and complications.
Results—Principal indications for upgrade were pacemaker syndrome (17), “opportunistic”—that is, at elective generator replacement (8), heart failure (7), non-specific breathlessness/fatigue (7), and neurally mediated syncope (3). Mean (SD) upgrade procedure duration (82.9 (32.6) minutes) significantly exceeded mean VVI implantation duration (42.9 (13.3) minutes) and mean DDD implantation duration (56.6 (22.7) minutes) (both p < 0.01). Complications included pneumothorax (1), ventricular arrhythmia requiring cardioversion (2), protracted procedure (10), atrial lead repositioning within six weeks (8), haematoma evacuation (1), superficial infection (1), and admission to hospital with chest pain (1); 20 patients (45%) suffered one or more complications including four of the eight who underwent opportunistic upgrade.
Conclusions—Pacemaker upgrade takes longer and has a higher complication rate than either single or dual chamber pacemaker implantation. This suggests that the procedure should be performed by an experienced operator, and should be undertaken only if a firm indication exists. Patients with atrial activity should not be offered single chamber ventricular systems in the belief that the unit can be upgraded later if necessary at minimal risk.

(Heart 1998;79:383–387)

Keywords: pacemaker upgrade; dual chamber pacemaker; single chamber pacemaker

Since the introduction of the dual chamber pacemaker, “upgrading” of single chamber ventricular pacemakers to dual chamber units has been a possibility in patients with persisting atrial electrical activity. Current guidelines suggest that patients with intrinsic atrial activity should receive dual chamber implants (or single chamber atrial implants if appropriate), but in the United Kingdom 40% of patients paced for high grade atrioventricular block during 1995 received fixed rate ventricular pacing systems (National Pacemaker Database, 1996) for reasons that were likely to include age, cost, relative inactivity, and intermittency of atrioventricular block. Patients receiving such relatively simple systems lack atrioventricular synchrony and a proportion (variously estimated at between 15% and 70%) go on to experience symptoms of the pacemaker syndrome. These patients, and those whose symptoms persist for other reasons (such as heart failure), may be considered for pacemaker upgrade.

There has only been one study concerning upgrading of pacemakers. This was undertaken in an asymptomatic population who underwent opportunistic upgrade at the time of elective generator change. There are no data relating to clinical experience of pacemaker upgrade in other patients. We sought to address this issue in a cardiology centre with a large pacemaker practice, assessing procedural characteristics, complication rates, and symptom response.

Methods
In this specialist cardiothoracic unit, approximately 350 permanent pacing systems are implanted annually. In the period under review (June 1989 to June 1997), 2614 patients received permanent pacemakers: 794 (31%) dual chamber, 1820 (69%) single chamber. Of those who received single chamber units, 986 were paced for high grade atrioventricular block (628) or sinus node disease (358), of whom 44 (4.5%) have subsequently undergone upgrading of single chamber ventricular units to dual chamber units. These patients formed the study group. Information regarding indications for upgrade, procedure duration, complications, symptom response, and subsequent clinical course were analysed. Complications of the pacing procedure were taken to include protracted upgrade procedure (defined as mean dual chamber implantation time plus twice the standard deviation from a random sample of 100 age matched implants, that is, > 102 minutes), recognised procedural adverse events (for example, pneumothorax, arrhythmia requiring cardioversion), pacemaker re-explosion within six weeks, or other hospital admission as a result of upgrade. These outcomes were recorded, and symptom response to upgrade was graded subjectively according to a three point scale.
Table 1  Original indication for pacing (n = 44)

<table>
<thead>
<tr>
<th>Indication</th>
<th>No of patients</th>
</tr>
</thead>
<tbody>
<tr>
<td>Complete heart block</td>
<td>17</td>
</tr>
<tr>
<td>Sinus node disease</td>
<td>8</td>
</tr>
<tr>
<td>Second degree atrioventricular block</td>
<td>7</td>
</tr>
<tr>
<td>Paroxysmal atrial fibrillation with pauses in ventricular response</td>
<td>3</td>
</tr>
<tr>
<td>Trifascicular block</td>
<td>3</td>
</tr>
<tr>
<td>Neuroly mediated syncope</td>
<td>3</td>
</tr>
<tr>
<td>Biphasic atral lead block with syncope</td>
<td>2</td>
</tr>
<tr>
<td>Hypertrophic cardiomyopathy</td>
<td>1</td>
</tr>
</tbody>
</table>

Table 2  Indication for upgrade of pacing unit and symptomatic response to upgrade

<table>
<thead>
<tr>
<th>Indication</th>
<th>No of patients (n = 44)</th>
<th>Symptom resolution following upgrade</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pacemaker syndrome</td>
<td>17</td>
<td>Complete: 11; Partial: 4; None: 2</td>
</tr>
<tr>
<td>Opportunistic (at generator change)</td>
<td>8</td>
<td>Not applicable*</td>
</tr>
<tr>
<td>Symptoms of heart failure</td>
<td>7</td>
<td>0; 3; 4; 3*</td>
</tr>
<tr>
<td>Non-specific breathlessness or fatigue</td>
<td>7</td>
<td>0; 4; 3; 3*</td>
</tr>
<tr>
<td>Neuroly mediated syncope</td>
<td>3</td>
<td>2; 1; 0</td>
</tr>
<tr>
<td>Progression to complete heart block</td>
<td>1</td>
<td>1; 0; 0</td>
</tr>
<tr>
<td>Prevention of paroxysms of atrial fibrillation</td>
<td>1</td>
<td>0; 0; 0</td>
</tr>
</tbody>
</table>

*One patient noted significant improvement in overall wellbeing.
†Includes one patient in whom the upgrade procedure could not be completed.

Table 3  Procedure duration

<table>
<thead>
<tr>
<th>Procedure</th>
<th>No of patients</th>
<th>Mean (SD) time (minutes)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Single chamber pacemaker implantation</td>
<td>100</td>
<td>42.9 (13.3)*</td>
</tr>
<tr>
<td>Dual chamber pacemaker implantation</td>
<td>100</td>
<td>56.6 (22.7)</td>
</tr>
<tr>
<td>Ventricular pacemaker upgrade</td>
<td>44</td>
<td>82.9 (32.6)*</td>
</tr>
</tbody>
</table>

*Single v dual p < 0.05; †single v upgrade p < 0.001; ‡dual v upgrade p < 0.001.

COMPLICATIONS
Ten patients underwent protracted procedures (defined above, > 102 minutes) Reasons included difficult dissection (3), adaptation of atrial or ventricular lead (2), difficulty establishing venous access (1), repeated atrial displacement (1), persistent left sided superior vena cava (1), difficulty in establishing adequate atrial parameters (1), and inability to sense or pace atrium (1).

Procedural adverse events were encountered in three patients. These were ventricular arrhythmia requiring DC cardioversion (2) and pneumothorax (1). Nine patients (20%) underwent re-exploration of their pacing system during the 7 days of the upgrade. Of these, eight were for atrial lead displacement (seven within 24 hours) and one for haematoma evacuation. Two patients were readmitted to hospital in the postoperative period for symptoms related to the upgrade procedure, one for superficial infection over the pacing generator (on three separate occasions), and one for chest pain. In total, 20 patients (45%) experienced one or more complications (fig 1). Median hospital stay was two nights, but eight patients (18%) remained in hospital for one or more extra nights as a result of complications.

UPGRADE AT ELECTIVE UNIT REPLACEMENT
Eight patients underwent “opportunistic” upgrade only, in that their pacing system was upgraded at the time of elective generator replacement despite the fact that they were asymptomatic in VVI mode. Of these patients, three had protracted procedures, one developing superficial infection requiring subsequent hospital admission. One further patient underwent repositioning of the atrial lead at two weeks. Therefore four of eight patients who underwent opportunistic upgrade of their pacing system encountered complications.

PROCEDURAL CHARACTERISTICS
The original implant was via the cephalic vein in seven patients and the subclavian vein in the remaining 37. Upgrade approaches were cephalic (4) and subclavian (40). In 11 patients the approach used for the upgrade was different from that used for the original implant; procedure duration (median (SEM)) did not differ according to whether the venous approach employed for the upgrade was different or the same (70 (12.6) v 80 (5.7) minutes, respectively, p = 0.21), nor did complication rates differ significantly.

The atrial lead initially chosen for use was passive fixation in 32 patients and active in 12. Of these, three passive atrial leads failed to

DATA ANALYSIS
Data distribution was assessed for normality. Data are expressed as mean (SD), or median (SEM), depending on distribution. Comparison of procedure duration was made with the Mann-Whitney U test, or the median test as appropriate. Comparison between procedure outcomes or characteristics was made with the χ² test or Fisher’s exact test as appropriate. The null hypothesis was rejected at the level p < 0.05.

Results
Between June 1989 and June 1997, 44 patients underwent attempted upgrade of a single chamber ventricular pacing system to a dual chamber system. Fifteen (36%) were female and 29 (64%) male. Average age at the time of original implant was 62.3 (13.8) years. Indications for original implant are shown in table 1.

Thirty three patients benefited symptomatically following original implant. Six required re-exploration of their original pacing system, for haematoma evacuation (1), ventricular lead repositioning (2), infection (1), lead erosion (1), and persistent syncope (1). Three patients underwent elective generator replacement in the interval between initial implant and upgrade.

The interval between original implant and upgrade was 5.9 (4.5) years, and the average age at upgrade was 68.2 (12.9) years. Indications for upgrade and symptom response to the procedure are shown in table 2. Of the 44 attempted upgrade procedures, 43 were successfully completed.

PROCEDURE DURATION
Upgrade procedure duration (82.9 (32.6) minutes) significantly exceeded single chamber pacemaker implantation (42.9 (13.3) minutes, p < 0.01) or dual chamber pacemaker implantation (56.6 (22.7) minutes, p < 0.01), both assessed from 100 randomly selected age matched procedures (table 3).
Protracted procedure or adverse procedural events

![Diagram](https://via.placeholder.com/150)

**Figure 1 Complications of 44 upgrade procedures.**

The most striking findings of our study concern procedure duration and early complications. Pacemaker upgrade took, on average, twice as long as single chamber ventricular implant, and almost half an hour longer than dual chamber implant, illustrating the technically more challenging nature of the procedure. Dissection of fibrosed and adherent tissue planes, attainment of venous access avoiding damage to the existing lead, and adaptation of ventricular or atrial leads to fit a new or old pacing systems at elective generator replacement. These patients experienced an improvement in perceived general wellbeing, and an increase in exercise capacity in DDD mode, but complications of the upgrade procedure which might have offset this potential benefit were not discussed. Other studies have compared VVI, DDI, and DDD modes, but these studies are confined to patients with dual chamber pacing systems already in situ.

The two patients subsequently underwent late pacing system revision, 15 and 48 months after upgrade respectively, both for ventricular lead failure. The upgrade procedure was not directly implicated in either case, though this possibility cannot be discounted. Seven patients have subsequently died, all of whom had poor left ventricular function at the time of upgrade, and four of whom underwent upgrade in an attempt to improve symptoms of heart failure. The interval between upgrade and death was 2, 4, 5, 6, 10, 28, and 30 months, respectively. All deaths were caused by heart failure, and none was attributable to the pacing procedure.

**Discussion**

Many patients undergo single chamber ventricular pacemaker implantation despite the presence of organised atrial electrical activity (UK National Pacemaker Database, 1996). Some of these patients later require system upgrade as a result of pacemaker syndrome or unresolved symptoms. Pacing upgrade is a more complex procedure than initial system implantation, but there are no studies assessing procedural characteristics and complications of this procedure. Sulke et al took 16 patients asymptomatic in VVI mode and upgraded their pacing systems at elective generator replacement. These patients experienced an improvement in perceived general wellbeing, and an increase in exercise capacity in DDD mode, but complications of the upgrade procedure which might have offset this potential benefit were not discussed. Other studies have compared VVI, DDI, and DDD modes, but these studies are confined to patients with dual chamber pacing systems already in situ.

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procedure, procedural complications, pacemaker re-exploration within six weeks, or other hospital admission as a result of upgrade were seen more commonly, though not to a level of statistical significance, among relatively inexperienced operators (<100 procedures). Complications were significantly more common, however, when comparing operators inexperienced with those experienced at upgrade procedures. Complications were statistically reduced after an operator’s second upgrade procedure, consistent with a steep but important learning curve.

Greatest symptomatic benefit was seen in those who underwent upgrade for pacemaker syndrome, neurally mediated syncope (with cardioinhibitory component), or development of complete heart block, as would be expected.11–15 18 19 Seven patients underwent pacemaker upgrade in an attempt to improve symptoms of heart failure, as DDD pacing in these patients has been shown to be of potential benefit.20–22 Results were relatively disappointing, in that four derived no symptomatic benefit while three benefited to some extent. Least clinical improvement was seen in those who underwent upgrade for non-specific breathlessness or fatigue in the absence of objective evidence of heart failure.

“Opportunistic” upgrading of a single chamber ventricular pacemaker to a dual chamber unit at the time of elective unit replacement remains contentious. Morbidity and mortality are greater in patients with persisting atrial activity paced in VVI mode than in modes which preserve atioventricular synchrony,17–20 but this does not necessarily mean that these patients will benefit from an upgrade procedure with its attendant risks. Sulke et al showed that patients in asymptomatic complete heart block can benefit from upgrade,11 and therefore suggested the concept of “subclinical” pacemaker syndrome, but potential complications of the procedure itself were not taken into account. In our series, eight patients underwent opportunistic upgrade, four of whom encountered problems. The potential benefit of upgrading in our asymptomatic subjects was therefore more than offset by the increased morbidity associated with the procedure itself. We suggest that pacemaker upgrade is not usually warranted in asymptomatic individuals, and should only be undertaken after careful consideration, by an operator experienced in upgrade procedures.

Pacemaker upgrade may be carried out with active or passive fixation atrial electrodes. It may be prudent to consider active fixation if the patient has previously undergone cardiac surgery,20 but this precaution is not always necessary, and may make use of the cephalic approach awkward. We found no significant difference in complication rate between passive and active atrial lead selection.

Mode of venous access for upgrade did not significantly influence complication rate, but the four patients who underwent cephalic upgrade following subclavian implant experienced no complications. The cephalic upgrade approach minimises risk of damage to the existing pacing lead, and removes risk of pneumothorax. We suggest that if the original implant was by subclavian vein puncture then upgrade should be undertaken through the cephalic vein.

The UKPACE trial will compare VVI, VVIR, and DDD pacing modes in elderly patients (over 70 years) with high grade atrioventricular block, and should provide clarification of the benefits (or otherwise) of sophisticated pacing in such patients. However, pending the results of this trial, the incidence of complications at upgrade that we have shown should mitigate against the use of VVI pacing in patients with ongoing atrial activity, a practice which remains commonplace despite the fact that the true incidence of pacemaker syndrome in such patients may exceed 50%.6 8 11 32 If expertise and budgets permit, these patients should be offered dual chamber units at the outset, rather than risk the need for upgrade at a later date.

CONCLUSIONS
Pacemaker upgrade is technically challenging and requires particular expertise. It should not be undertaken in asymptomatic subjects without considerable thought, and should be performed by an experienced operator. Pacemaker upgrade takes longer and has a higher complication rate than either single or dual chamber pacemaker implantation. Patients with atrial activity should not be offered single chamber ventricular systems on the pretext that the unit can be upgraded if necessary at minimal risk.


