Pacing-associated cardiomyopathy in adult congenital heart disease

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ABSTRACT

Objectives Long-term single-site ventricular pacing may adversely affect ventricular function, due to dyssynchronous systemic ventricular contraction. We sought to determine the incidence, predictors and outcomes of pacing-associated cardiomyopathy (PACM) in an adult congenital heart disease (ACHD) cohort.

Methods We retrospectively identified all patients in our database with a ventricular pacemaker from 2000 to 2019. Patients were followed for the primary endpoint of unexplained decline in systemic ventricular function (PACM) and the secondary endpoint of heart failure admission.

Results Of 2073 patients in our database, 106 had undergone pacemaker implantation. Over a median follow-up of 9.4 years, 25 patients (24%) developed PACM, but only in those with ventricular pacing percentage (VP%) ≥70%; PACM occurred in 0% of those with VP <70% and 47% of those with VP ≥70% (p=0.001). High-burden ventricular pacing (≥70%) remained predictive of PACM in transposition of the great arteries, tetralogy of Fallot and complex biventricular repair subgroups, but not in Fontan patients. Those with PACM were more likely to be admitted with heart failure (44% vs 15%, p=0.002). Cardiac resynchronisation therapy (CRT) upgrade was performed in 11 patients, with 9 responders (82%).

Conclusions In a cohort of patients with ACHD followed long-term post-pacing, 24% developed cardiomyopathy that was significantly associated with a higher burden of ventricular pacing (VP ≥70%). Given promising response rates to CRT, patients with ACHD expected to pace in the ventricle should be closely monitored for systemic ventricular decline.

INTRODUCTION

The potential adverse effects of chronic single-site ventricular pacing have been well described in acquired heart disease.1,2 Persistent dyssynchronous left ventricular (LV) contraction, due to right ventricular pacing, may result in systolic LV impairment.3 This is termed ‘pacing-induced cardiomyopathy’, in the absence of another cause.3,4 The risk of clinical heart failure is also increased in these individuals.1,2 At a histopathological level, myocardial biopsies from young patients with congenital complete heart block (CHB) and chronic pacing have demonstrated degenerative LV fibrosis.5,6 The prevalence of significant bradycardia requiring pacing in patients with adult congenital heart disease (ACHD) substantially exceeds the age-equivalent general population.7 Pacing-makers are most commonly dual chamber, with single-site endocardial or epicardial ventricular leads. Propensity to ventricular pacing depends on the initial indication for implant; bradycardia in ACHD may result from surgical trauma and/or scarring to the sinus or atrioventricular (AV) node, progressive atrial fibrosis, treatments for atrial arrhythmia or an anatomical abnormality.
of the conduction system. Concerningly, patients with ACHD may be more vulnerable to the adverse effects of ventricular pacing, since they also have multiple predisposing risk factors for heart failure. Certain congenital lesions, such as those with a systemic right ventricle, have a predisposition to ventricular failure; differentiating this from pacing-induced cardiomyopathy is challenging. Nevertheless, identification of ‘at-risk’ individuals is valuable, as cardiac resynchronisation therapy (CRT) is an effective intervention in both acquired heart disease and ACHD. We sought to determine the incidence, predictors and outcomes of pacing-associated cardiomyopathy in a population with ACHD.

METHODS
Study population and clinical characteristics
We retrospectively identified all patients with ACHD in our database with a permanent pacemaker (PPM), who had been seen at least two times in our unit between 2000 and 2019. Our database represents a quaternary referral centre with active follow-up of over 2000 patients with ACHD. Patients with congenital CHB and a structurally normal heart were excluded, as were patients with an implantable defibrillator or a de novo CRT implant. Clinical history was extracted from the medical records, with investigations taken from the most recent visit prior to PPM implantation. Systemic (subaortic) ventricular systolic function was measured using Simpson’s rule for subaortic left ventricles and subjective grading for subaortic right ventricles. Mild dysfunction was defined as an ejection fraction of 40%–49%, moderate 30%–39% and severe <30%. Complexity of CHD was classified according to the Bethesda criteria. Patients were not involved in this research.

Pacing-induced cardiomyopathy
The indication for PPM implant was categorised as sinus node disease, AV block, post-surgical sinus node disease, post-surgical AV block, tachy–brady syndrome or other. Patients were then followed for the primary endpoint of pacing-associated cardiomyopathy (PACM). This was defined as a decline in ejection fraction of at least one grade (≥10% if subaortic LV, or subjective assessment if subaortic right ventricle), to a value <50%, in the absence of another identifiable aetiology (eg, valvular heart disease). Given patients with ACHD may develop ventricular impairment due to the natural progression of their disease, the ‘PACM’ endpoint was comprised of cardiomyopathy deemed ‘possibly’ or ‘probably’ related to ventricular pacing, as assessed by an ACHD cardiologist and electrophysiologist. The secondary endpoint was heart failure admission for any cause. Cardiomyopathy was chosen as the primary endpoint, rather than heart failure admission, as these were not patients with heart failure per se, and PACM may develop in the absence of symptoms; ventricular impairment is associated with adverse clinical outcomes in the setting of complex CHD. Prespecified subanalysis was performed stratifying patients into two groups based on age at PPM implant: PPM implant <18 years old vs ≥18 years old. This cut-off was chosen to further assess long-term outcomes of paediatric versus adult pacing. Ventricular pacing percentage (VP%) was recorded at first review post-PPM implant and at last review. VP% at last review was recorded immediately prior to the development of PACM, or at last follow-up for those who did not develop this endpoint. VP% at last review was used for the purposes of predictive analysis. In the patients who underwent upgrade to CRT, a ‘non-responder’ was defined as those with no improvement in ejection fraction, or a decline in ejection fraction, post-CRT.

Statistical analysis
Continuous variables are presented as mean±SD, or median±IQR. Categorical variables are presented as frequencies with percentages. A receiver-operating characteristic (ROC) curve was constructed to assess the optimal cut-off for VP% that best predicted the PACM endpoint. Survival free from PACM was assessed by the Kaplan-Meier method. Variables associated with PACM or heart failure admission were assessed by the X², Fisher exact or t-test as appropriate, on univariate analysis. Variables assessed were as follows: age, gender, systemic right ventricle, complex CHD, prior heart failure episode, QRS width, QRS morphology (left bundle branch block), pre-existing systemic ventricular impairment and VP%. Multivariate logistic regression models were performed to identify independent predictors of PACM, incorporating the following variables added sequentially: demographics (age and gender), systemic ventricular morphology (left, right or univentricular), VP%, duration of pacing. A two-sided p value of <0.05 was considered statistically significant. Statistical analysis was performed using SPSS V.22.0.

RESULTS
Study population
Of 2073 patients in our database, we identified 106 patients with a PPM, who were seen at least two times in our unit between 2000 and 2019 (5.1%). The median age at implant was 29 years (IQR 16–40), with a median follow-up post-implant of 9.4 years (IQR 4.0–21.5). Thirty-four patients were less than 18 years old at time of implant (32%; median age 9), and 72 patients were greater than 18 years old (68%; median age 36). CHD diagnoses are displayed in figure 1. Baseline characteristics of the PPM patients are shown in table 1. Of 55 patients with a history of supraventricular arrhythmia, 35% had previously undergone catheter ablation; the AV node was intentionally ablated in one patient.

Indications and implants
PPM implantation was for sinus node disease in 29 (27%), tachy–brady syndrome in 13 (12%), AV block in 34 (32%), post-surgical sinus node disease in 3 (3%), post-surgical AV block in 20 (19%) and other reasons

(for example, syncope with suspected bradycardia) in 7 (7%). The total proportion of pacemakers implanted for AV block was 51%. Devices were dual chamber in 71 patients (67%), atrial lead only in 3 (3%) and ventricular lead only in 31 (30%). Ten of the single chamber devices were subsequently upgraded to dual chamber (one atrial, nine ventricular). Atrial leads were endocardial in 65% and epicardial in 35%. Ventricular leads were endocardial in 51% and epicardial in 49%. Early complications (within 30 days of implant) occurred in 13 patients (12%) and late complications in 29 patients (27%). Early complications included lead revision in five patients, pocket haematoma in three, pneumothorax in three, tamponade in one and early death in one. Late complications included lead revision/failure in 29 patients, endocarditis in 3 and allergy to device component in 1 (noting several patients suffered more than one complication). Mean VP% at first review post-implant was 51%±43% and at last review was 54%±44%; these were highly correlated (p<0.001). In those paced for ‘AV block’, mean VP% was 83%, and in all others mean VP% was 25% (p<0.001).

Pacing-associated cardiomyopathy

The primary endpoint of PACM occurred in 25 patients (24%), at a median time of 11.7 years post-implant (IQR 2.5–23.1). PACM was deemed ‘probably’ due to ventricular pacing in 19 patients and ‘possibly’ related in 6 patients. The primary endpoint, as a proportion of each CHD diagnosis, occurred as follows: transposition of the great arteries (TGA—atrial switch and congenitally corrected) in 10/31 patients (32%), Fontan circulation in 2/24 patients (8%), other complex CHD in 4/12 (33%), tetralogy of Fallot (TOF) in 5/16 (31%) and other simple–moderate CHD in 4/23 (17%). PACM occurred in 10/34 patients with PPM implant at age <18 years old (29%) and 15/72 patients with PPM implant at age ≥18 years old (21%). The mean time to PACM was significantly longer in those paced from age <18 years old versus those paced aged ≥18 years old (mean 24±12 years vs 4±4 years post-implant, respectively, p<0.001). Congenital diagnoses were evenly distributed between the two age groups. A deterioration in ejection fraction with a defined cause, such as valvular heart disease, occurred in 7/81 patients not meeting the primary endpoint (9%). Heart failure admission for any cause occurred in 24 patients (23%), at a median time of 5.7 years post-implant. Patients with PACM were more likely to be admitted with heart failure, compared with those who did not develop PACM (44% vs 15%, p=0.002).

Predictors of PACM: ventricular-pacing burden

On univariate analysis, higher VP% alone was predictive of PACM (p<0.001), as a continuous variable. Specifically,
PACM in those <18 years old at time of PPM implant and was more often paced for sick sinus syndrome, with a smaller proportion pacing ≥70% in the ventricle. When PACM ≥70% remained significantly associated with a high burden of ventricular pacing (VP ≥ 70%), with surgical epicardial leads in four patients (TGA post-Mustard, congenitally corrected TGA, Ebstein’s anomaly, aortic stenosis). A total of 11 patients from the entire cohort were upgraded to an implantable cardioverter-defibrillator, without CRT capability (10%). There were 14 deaths (13%) during follow-up, at a mean time of 11.4 years post-PPM implant (range 0.1–27.8 years). Deaths were due to heart failure (n=8), infective endocarditis (n=2), pulmonary hypertension/respiratory failure (n=2) and cerebrovascular events (n=2). In the group who met the primary endpoint, four deaths occurred: one due to stroke and three due to heart failure (one Mustard patient who failed to respond and one who died from recurrent hospitalisation). The interventions and outcomes in the 25 patients who met the primary endpoint are shown in table S3. A total of 11 patients were upgraded to CRT pacemaker±defibrillator, with 9 responders and 2 non-responders (ie, responder rate 82%). CRT upgrade is planned, but not yet performed, in a further three patients. In the CRT upgrades, systemic ventricular leads were implanted endocardially via the coronary sinus in six patients (four TOF, one congenitally corrected TGA, one repaired truncus arteriosus), endocardially via a persistent left-sided superior vena cava in one patient (repaired ventricular septal defect), with surgical epicardial leads in four patients (TGA post-Mustard, congenitally corrected TGA, Ebstein’s anomaly, aortic stenosis). A total of 11 patients from the entire cohort were upgraded to an implantable cardioverter-defibrillator, without CRT capability (10%). There were 14 deaths (13%) during follow-up, at a mean time of 11.4 years post-PPM implant (range 0.1–27.8 years). Deaths were due to heart failure (n=8), infective endocarditis (n=2), pulmonary hypertension/respiratory failure (n=2) and cerebrovascular events (n=2). In the group who met the primary endpoint, four deaths occurred: one due to stroke and three due to heart failure (one Mustard patient who failed to respond to CRT, as well as Fontan and Rastelli patients who were too unwell for intervention).

**DISCUSSION**

In a cohort of patients with ACHD followed long-term post-pacemaker implantation, one in four developed an otherwise unexplained decline in systemic ventricular function, significantly associated with a high burden of ventricular pacing (VP ≥ 70%). The adverse effects of long-term single-site ventricular pacing have been appreciated for some time in the population with acquired heart disease. Ventricular pacing results in electrical and mechanical dyssynchrony of the systemic ventricle, reducing effective stroke volume. At a histopathological level, myofibrillar disarray has been observed in a canine-pacing model and degenerative fibrosis in biopsies from chronically paced young patients with congenital CHD. Despite this, a subset of patients appear to tolerate long-term ventricular pacing well; risk factors for pacing-induced cardiomyopathy in acquired heart disease and those ≥18 years old, analysed independently. Post-hoc analysis excluding all ‘possible’ patients with PACM (ie, including only ‘probable’ PACM) did not change the significant association of VP ≥ 70% with PACM. Ventricular lead pacing site did not significantly affect rates of PACM (15/53 endocardal vs 10/51 epicardial leads developed PACM; 28% vs 20%; p=0.300). There was no significant difference in mean VP% between those patients with endocardial versus epicardial leads (55% vs 48%; p=0.154). There were no significant predictors of ‘heart failure admission for any cause’.

**Outcomes in patients with PACM**

<table>
<thead>
<tr>
<th>CHD lesion</th>
<th>Mean VP (%)</th>
<th>PACM endpoint</th>
<th>VP &lt;70%</th>
<th>VP ≥70%</th>
<th>P value</th>
</tr>
</thead>
<tbody>
<tr>
<td>TOF</td>
<td>58</td>
<td>0/7 (0%)</td>
<td>5/9 (56%)</td>
<td>0.034</td>
<td></td>
</tr>
<tr>
<td>TGA*</td>
<td>57</td>
<td>0/14 (0%)</td>
<td>10/17 (59%)</td>
<td>0.004</td>
<td></td>
</tr>
<tr>
<td>Fontan</td>
<td>34</td>
<td>0/17 (0%)</td>
<td>2/7 (29%)</td>
<td>0.130</td>
<td></td>
</tr>
<tr>
<td>Complex repair</td>
<td>62</td>
<td>0/3 (0%)</td>
<td>4/5 (80%)</td>
<td>0.048</td>
<td></td>
</tr>
</tbody>
</table>

*Atrial switch and congenitally corrected, that is, those with systemic right ventricles.
CHD, congenital heart disease; TGA, transposition of the great arteries; TOF, tetralogy of Fallot.

Figure 2 Survival free from pacing-associated cardiomyopathy (PACM). PPM, permanent pacemaker; VP, ventricular pacing.
Congenital heart disease

Table 3  Interventions and outcomes in patients who developed pacing-associated cardiomyopathy

<table>
<thead>
<tr>
<th>CHD diagnosis</th>
<th>N</th>
<th>Intervention/outcome</th>
<th>Response to CRT</th>
</tr>
</thead>
<tbody>
<tr>
<td>TOF</td>
<td>5</td>
<td>CRT upgrade (n=3)</td>
<td>3/3 responded</td>
</tr>
<tr>
<td></td>
<td></td>
<td>CRT upgrade intended (n=1)</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Lost to follow-up (n=1)</td>
<td></td>
</tr>
<tr>
<td>TGA*</td>
<td>10</td>
<td>CRT upgrade (n=4)</td>
<td>3/4 responded</td>
</tr>
<tr>
<td></td>
<td></td>
<td>CRT upgrade intended (n=1)</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Medical therapy only (n=2)</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Listed for heart/lung transplant (n=1)</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Death (n=1)</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Lost to follow-up (n=1)</td>
<td></td>
</tr>
<tr>
<td>Fontan</td>
<td>2</td>
<td>Medical therapy only (n=1)</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Death (n=1)</td>
<td></td>
</tr>
<tr>
<td>Other complex CHD</td>
<td>4</td>
<td>CRT upgrade (n=1)</td>
<td>1/1 responded</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Improved with reduction in VP% (n=1)</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Medical therapy only (n=2)</td>
<td></td>
</tr>
<tr>
<td>Other simple–moderate CHD</td>
<td>4</td>
<td>CRT upgrade (n=3)</td>
<td>2/3 responded</td>
</tr>
<tr>
<td></td>
<td></td>
<td>CRT upgrade intended (n=1)</td>
<td></td>
</tr>
</tbody>
</table>

*Atrial switch and congenitally corrected, that is, those with systemic right ventricles.

CHD, congenital heart disease; CRT, cardiac resynchronisation therapy; TGA, transposition of the great arteries; TOF, tetralogy of Fallot; VP%, ventricular pacing percentage.

include pre-existing systemic ventricular impairment and a higher burden of ventricular pacing. Patients with ACHD may be especially susceptible to this phenomenon, as this population has multiple predisposing risk factors for heart failure, depending on the underlying lesion.

In acquired heart disease, approximately 10%–20% of patients will develop systemic ventricular dysfunction when followed long-term post-pacing. Our incidence of 24% exceeds these reports, but may overestimate true ‘pacing-induced cardiomyopathy’ due to the natural history of systemic ventricular decline in certain congenital lesions, such as those with systemic right ventricles. In patients with ACHD it is inherently difficult to delineate the relative contribution of pacing from natural disease progression particularly in those with pre-existing ventricular impairment prior to pacing, or from cardiomyopathy related to surgical intervention. Interestingly, in our study those patients paced in adulthood typically developed PACM in the short midterm (mean 4 years) and those paced in childhood typically developed PACM very late post-implant (mean 24 years). This may suggest that a significant proportion of patients with ACHD paced in childhood developed ventricular decline due to natural disease progression, rather than pacing. Pacing-induced cardiomyopathy in acquired heart disease cohorts usually occurs in the first 4 years, however, in long-term follow-up of children paced for congenital CHB, up to 10% developed LV impairment at a mean time of 15 years post-implant. It is possible that a minority of vulnerable-paced young patients may suffer very late cardiomyopathy, despite many years of stable ventricular function and thus lifelong monitoring of ventricular function is warranted. Pacing dyssynchrony may also exacerbate late systemic ventricular decline due to natural disease progression in ACHD. Incidence of PACM in ‘whole of ACHD’ cohorts has not previously been assessed. The relatively low rates of systemic ventricular decline due to a specific (non-PACM) aetiology may reflect the young average age of our population.

The development of PACM was significantly associated with a high burden of ventricular pacing (VP ≥70%) in our study. This ‘exposure–response’ relationship further supports the pathophysiological link between chronic dyssynchrony and ventricular impairment. Importantly, this association does not prove causation, as patients with more extensive myocardial and/or electrical disease are more likely to require pacing. Previously, small single-lesion studies in ACHD have supported a correlation between moderate-high burden ventricular pacing and cardiomyopathy in univentricular hearts and congenitally corrected TGA, but failed to show this in TOF. In our study, the correlation between high-burden pacing and cardiomyopathy was consistent across TGA (atrial switch and congenitally corrected), TOF and complex biventricular repair subgroups. Conversely, most Fontan patients were paced for sick sinus syndrome and thus both high-burden ventricular pacing and PACM were uncommon. The preferred ‘cut-off’ in VP% that best predicts cardiomyopathy is contentious. In the MOST trial, a VP >40% was predictive of heart failure hospitalisation. This cut-off was then adopted and supported by several subsequent studies, although cardiomyopathy with VP >20% has also clearly been described. The proportion of patients with ventricular pacing between 20% and 70% in our study...
was relatively small, and so our study may not have been powered to detect this association. We suggest vigilance is required in any patient with ACHD expected to have even a modest burden of ventricular pacing. Notably, in contrast to acquired heart disease, pre-existing systemic ventricular impairment was not associated with the development of PACM. Those patients who developed PACM were more likely to be admitted with heart failure in our study, but VP% per se was not predictive of ‘heart failure admission’. This may be due to the multiple heterogeneous aetiologies driving heart failure admission in ACHD, diluting the significance of ventricular pacing alone.

In patients exposed to unavoidable pacing with declining systemic ventricular function, upgrade to CRT should be considered. In those who were upgraded to CRT in our study, response rates were high, exceeding 80%. This supports the ACHD literature, which has reported CRT response rates of 80%–90%, favourable in comparison with acquired heart disease. Despite the anatomical complexities in ACHD, complication rates of CRT implant or upgrade were not prohibitive in these studies. The subgroup of patients with ACHD with PACM and upgrade may have a greater likelihood of response than de novo CRT implants, highlighting the importance of making this diagnosis. Nevertheless, uncertainties do remain regarding CRT response rates within select CHD lesions. It is unclear if multisite algorithms to reduce VP% may be beneficial. In acquired heart disease, guidelines suggest this should be considered in patients with heart failure and reduced ejection fraction, with an expected high burden of ventricular pacing (typically >40%). This recommendation has been extrapolated to ACHD guidelines.

Limitations

There are several limitations to our study. Our study is retrospective and single-centre. Prevalence of sinus node disease versus AV block varies across differential congenital lesions, thus skewing the proportion of CHD diagnoses in the VP ≥70% vs VP <70% subgroups. Despite this, VP ≥70% remained significantly associated with PACM across multiple congenital lesions. Relative numbers of patients with VP% between 20% and 70% were low, and so this study may not have been powered to detect an association with PACM in this moderate pacing group. The use of AV nodal blocking or antiarrhythmic drugs was not analysed, but these may affect VP%. Differentiating pacing-induced cardiomyopathy from natural progression of disease is inherently difficult in ACHD, and thus our findings do not represent a true incidence of ‘pacing-induced cardiomyopathy’, but rather an association between ventricular pacing and the endpoint of deteriorating ventricular function. Clinical assessment is likely to overestimate ‘true’ pacing-induced cardiomyopathy incidence.

CONCLUSIONS

In a cohort of patients with ACHD followed long-term post-pacemaker implantation, a high burden of ventricular pacing (VP ≥70%) was significantly associated with the development of cardiomyopathy. This association was independently present across TGA, TOF and complex biventricular repair lesion subgroups. Patients with ACHD expected to pace in the ventricle should be closely monitored for PACM, and affected patients be considered for upgrading to biventricular pacing.

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 Contributors BM was responsible for research design, acquisition and interpretation of data, and critically revising the manuscript and approval of the final version. BM is responsible for the overall content as a guarantor. CM was responsible for research design, critically revising the manuscript and approval of the final version. MM was responsible for research design, critically revising the manuscript and approval of the final version. DSC was responsible for research design, critically revising the manuscript and approval of the final version. RLC was supported by a Health Professional Scholarship from the National Heart Foundation of Australia. BM was responsible for research design, acquisition and interpretation of data, and critically revising the manuscript and approval of the final version. DSC was responsible for the overall content as a guarantor. MM was responsible for research design, critically revising the manuscript and approval of the final version. RLC was responsible for research design, critically revising the manuscript and approval of the final version.

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