Early performance of a miniaturized leadless cardiac pacemaker: the Micra Transcatheter Pacing Study

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Received 9 February 2015; revised 30 March 2015; accepted 4 May 2015; online publish-ahead-of-print 4 June 2015

See page 2520 for the editorial comment on this article (doi:10.1093/eurheartj/ehv261)

Aims

Permanent cardiac pacing is the only effective treatment for symptomatic bradycardia, but complications associated with conventional transvenous pacing systems are commonly related to the pacing lead and pocket. We describe the early performance of a novel self-contained miniaturized pacemaker.

Methods and results

Patients having Class I or II indication for VVI pacing underwent implantation of a Micra transcatheter pacing system, from the femoral vein and fixated in the right ventricle using four protractible nitinol tines. Prespecified objectives were >85% freedom from unanticipated serious adverse device events (safety) and <2 V 3-month mean pacing capture threshold at 0.24 ms pulse width (efficacy). Patients were implanted (n = 140) from 23 centres in 11 countries (61% male, age 77.0 ± 10.2 years) for atrioventricular block (66%) or sinus node dysfunction (29%) indications. During mean follow-up of 1.9 ± 1.8 months, the safety endpoint was met with no unanticipated serious adverse device events. Thirty adverse events related to the system or procedure occurred, mostly due to transient dysrhythmias or femoral access complications. One pericardial effusion without tamponade occurred after 18 device deployments. In 60 patients followed to 3 months, mean pacing threshold was 0.51 ± 0.22 V, and no threshold was ≥2 V, meeting the efficacy endpoint (P < 0.001). Average R-wave was 16.1 ± 5.2 mV and impedance was 650.7 ± 130 ohms.

Conclusion

Early assessment shows the transcatheter pacemaker can safely and effectively be applied. Long-term safety and benefit of the pacemaker will further be evaluated in the trial.

Clinical Trial Registration

ClinicalTrials.gov ID NCT02004873.

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**Introduction**

Permanent cardiac pacing, the only effective treatment for symptomatic bradycardia, reduces symptoms and recurrence of syncope, and improves survival in high-risk populations.\(^1\)\(^-\)\(^3\) Since their introduction in the 1960s, pacemakers have shrunk in size and grown in sophistication, yet their basic function remains to be sustaining a normal heart rate. Conventional pacing systems consist of a pacemaker containing the electronics and battery typically implanted in a subcutaneous pocket in the chest. One or more leads threaded from the device pocket through veins into the heart conduct the pacing therapy to the desired pacing site. When veins cannot be used, a surgical procedure includes implanting epicardial leads that link the heart to the device. Despite the reduction in complications due to technological advances, serious adverse events can still be encountered. These are reported to be 20% at 5 years, with highest contributions related to the pacing lead (\(\sim\)11%) and pocket (\(\sim\)8%),\(^4\) including pneumothorax after subclavian vein puncture, pocket haematoma, erosion or infection, vein stenosis or occlusion, endocarditis, tricuspid valve trauma, connection troubles, lead fractures, and other malfunctions.\(^5\)\(^-\)\(^8\) Micra\(^\text{TM}\) transcatheter pacing system (TPS, Model MC1VR01, Medtronic plc, Mounds View, MN, USA) is a miniaturized single-chamber pacemaker system that is delivered via catheter through the femoral vein and implanted directly inside the right ventricle (RV) of the heart. This new technology is feasible due to advances in miniaturization (high-density battery), low-power electronics, catheter delivery systems, novel materials (nitinol), and electrodes being directly placed on the pacemaker capsule. This eliminates the need for a device pocket and insertion of a pacing lead, thereby eliminating an important source of complications associated with traditional pacing systems while providing similar benefits.

The purpose of this report is to summarize the early performance of the TPS using pre-specified safety and efficacy objectives from the Micra Transcatheter Pacing System Study. These data are being used to support European regulatory submission.

**Methods**

**Study design and oversight**

The trial is a prospective, multi-site, single-arm, worldwide clinical study evaluating the safety and efficacy of the TPS. Briefly, the study will implant up to 720 patients at up to 70 centres worldwide. All participants must satisfy standard criteria for de novo pacemaker implantation with Class I or II indications and provide written informed consent. The trial design is described in detail elsewhere.\(^9\) The trial is sponsored by the manufacturer, Medtronic. The protocol was approved by the Ethics Committee at each participating institution and associated national and local regulatory agencies. Data of this ongoing study are collected by trained centre personnel, and data integrity maintained via programmatic edit checks and source data verification by the sponsor. Safety and trial conduct oversight are provided by an independent data monitoring committee.

**Study device and implant procedure**

The transcatheter pacing system is a 0.8 cc, 2.0 g, self-contained, hermetically enclosed capsule, single-chamber ventricular pacemaker with functionality and features similar to existing ventricular pacemakers, inclusive of rate responsive pacing and automated pacing capture threshold management. The device is 25.9 mm in length, with an outer diameter of 6.7 mm. A nominal pulse width duration of 0.24 ms was chosen since it is near the chronaxie of the strength duration curve.\(^10\) Utilizing a pulse width near the chronaxie should minimize pacing energy and improve battery longevity. By design, it is conditionally safe for full body magnetic resonance imaging in 1.5 and 3.0 Tesla scanners. The device is fixated via four electrically inactive protractible nitinol tines located on the distal end of the device (Figure 1).

The TPS is tethered to and sits in a cup at the distal end of a steerable transfemoral catheter delivery system and placed through the femoral vein using a 23 French internal diameter/27 French outer diameter introducer (Figure 2). The introducer is advanced using a guide-wire and dilator into the right atrium. The guide-wire and dilator are then removed, and the steerable delivery system catheter with TPS preloaded and tethered is then advanced into the RV. Transcatheter pacing system

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**Figure 1** Transcatheter pacing system single-chamber ventricular pacemaker. Illustration of transcatheter pacing system positioned in the RV apex. RV, right ventricle.
is deployed by retraction of the device-containing cup at the distal end of the delivery catheter positioned against the RV endocardium with fixation into the myocardium by the associated protraction of the nitinol tines (Figure 3). The delivery catheter is then withdrawn several centimetres and the fixation confirmed by a ‘pull and hold’ test. The tether is pulled until counter movement of the heartbeats can be felt and the tine deflection can be observed fluoroscopically, with the tethering material still connected to the TPS. Although bench testing has shown a single tine engaged in tissue holds the device securely within the myocardium with a high margin of safety, it is recommended that two tines are engaged for further security of fixation. Therefore, investigators are requested to check fluoroscopically that at least two tines are engaged within the myocardium before releasing the device; otherwise, the device should be retracted and repositioned to another position within the RV. Once the device is placed in the RV and adequate device fixation verified, electrical measurements (pacing thresholds, pacing impedance, R-wave amplitude) are checked. Inadequate fixation (e.g. device tines engage into the trabeculae rather than RV myocardial wall) will be apparent via unacceptable electrical measurements and possibly failed pull and hold test. In case of persistence of high thresholds after two to three deployments, investigators were advised to remove the transcatheter delivery system in order to check for thrombus covering the tip electrode, and, if found, to carefully remove the thrombus. No specific implant location is recommended as the device can be placed at various anatomical RV positions. However, it is suggested to avoid placement at the free wall to minimize risk of effusion.

After adequate electrical measurements are obtained, the tether is cut and the delivery system is removed. Haemostasis is achieved via various closure methods, determined by implanter. Post-procedural haemostasis and peri-procedural antibiotics and anticoagulation are at implanter discretion with the exception that intra-procedural heparinized flushing of the introducer is recommended in all patients.

Therapy initiation and follow-up

Enrolled patients undergo system implant attempt and then are followed, including adverse event and device evaluation, at implant, hospital discharge, 1, 3, 6, and 12 months post-implant. Implanted patients are evaluated semi-annually until trial closure. Cardiovascular, procedure, and system-related adverse events are adjudicated by an independent clinical events committee.

Initially, pacemaker-dependent patients were excluded. Twenty-five implanted patients underwent 24 h ambulatory ECG and device function (markers and EGMs) monitoring (Model ER220, Medtronic plc, Mounds View, MN, USA). After comprehensive review of the ambulatory ECG and safety data by the trial’s independent data monitoring committee, allowance of pacemaker-dependent patients was determined.

Early performance objectives

Early performance objectives, the subject of this report, were assessed once the 60th patient completed the 3-month post-implant visit. At that point, 140 patients had been implanted (Figure 4). The safety objective was assessed in all 140 implanted patients, and the efficacy objective was assessed in the 60 subjects who had been followed through 3 months. These early performance objectives were required to obtain CE mark:

(i) The early safety performance objective was to demonstrate that the freedom from unanticipated serious adverse device events (USADEs) was significantly >85% in all implanted patients once 60 patients were followed through 3 months. The early safety performance objective was assessed in all 140 implanted patients. USADEs are defined as serious adverse events related to the use of the TPS not previously identified in nature, severity, or degree of incidence. USADEs were selected for the early safety assessment endpoint since the risk profile of transvenous pacemaker lead systems has been well-established. Thus, the rate of USADEs was selected to characterize unforeseen risk associated with TPS.

(ii) The early efficacy performance objective was to demonstrate that the mean pacing capture threshold at pulse duration of 0.24 ms was significantly lower than 2 V in the 60 patients who completed the 3-month follow-up visit. Meeting this objective ensures TPS would have battery longevity estimation as expected.
In addition to the USADE early performance objective, all adverse events from the 140 implanted patients were carefully described and reviewed by the Data Monitoring Committee.

**Statistical methods**

A sample size of 60 patients successfully implanted with TPS and followed for at least 3-months post-implant provided more than 90% power at a type I error level of 0.025 to test the null hypothesis associated with the early performance efficacy objective assuming the true mean 3-month pacing capture thresholds is 1.0 V with a standard deviation of 0.5 V. This sample size provided more than 90% power for testing the early performance safety objective since all patients with an implant attempt at the time the 60th 3-month visit was accrued would be included in the analysis of the safety objective.

The Kaplan–Meier method was pre-specified as the method for evaluating the safety objective so all patients with an implant attempt could be included in the analysis. However, since no USADEs were observed, the exact binomial test comparing the observed 3-month USADE free rate to the null value of 85% was used to evaluate the safety objective and derive the lower 97.5% confidence interval (CI). A one-sample t-test comparing the observed mean 3-month pacing capture threshold to the null value of 2.0 V was used to test the efficacy objective. In addition, paired t-tests were used to compare electrical variables measured at implant and the 3-month visit. Statistical calculations were performed using SAS (SAS Institute, Cary, NC, USA) or R ([www.r-project.org](http://www.r-project.org)) and validated per the sponsor’s operating procedures. The procedure duration was defined as the time from the insertion of the TPS introducer to removal. Time to hospital discharge was defined as the number of days from implant to hospital discharge.

**Results**

Patient recruitment began 5 December 2013 and the 60th 3-month visit accrued on 11 August 2014, triggering evaluation of the early performance objectives. At the time of database closure for this analysis, 140 patients had an attempted implant and all were successfully implanted. The TPS was implanted in these 140 patients in 23 study centres by 37 physicians in 11 countries and were followed to an average of 1.9 ± 1.8 months (range 0–6.5 months).
Implant attempted \( (n = 140) \)

| Received TPS \( (n = 140) \) | Did not receive TPS \( (n = 0) \) |

Follow-up

| Death \( (n = 1) \) | Lost to follow-up \( (n = 0) \) | Discontinued intervention \( (n = 0) \) |

Analysed for early performance objectives

| Safety \( (n = 140) \) | Three-month efficacy \( (n = 60) \) |

Figure 4  Flow diagram of patients analysed. Flow diagram from patients implanted by 11 August 2014 and analysed for early performance objectives.

Patients were mostly male (60.7%), of mean age 77.0 ± 10.2 years, height range 144–190 cm, and weight range 41–148 kg (Table 1). The most common primary indications for pacing were atrioventricular (AV) block (66%, or 93 of 140 patients) and symptomatic sinus node dysfunction (29%, or 40 of 140 patients). Nine (6%) patients were not felt to be appropriate for implantation of conventional transvenous lead pacing systems due to a variety of reasons such as compromised venous access, previous infection, and cancer with need for indwelling catheter. Confirmation of appropriate pacemaker operation from the 24 h ambulatory ECG analysis by the trial’s data monitoring committee was achieved 3 weeks prior to database closure, enabling pacemaker-dependent patient enrolment. Two pacemaker-dependent patients were subsequently implanted and included in this analysis.

Ninety-one (65%) of the 140 patients received a VVI pacemaker for bradycardia in conjunction with permanent or persistent atrial tachyarrhythmias. Of the remaining 49 patients, 22 had sinus node dysfunction, 19 had AV block, two had sinus node dysfunction plus AV block, and six had other reasons listed for choosing a ventricular pacemaker. In these 49 patients, the predominant reason for TPS selection was identified as ‘infrequent pacing expected’ (69%) and ‘advanced age’ (22%). Other reasons included sedentary lifestyle, anatomical limitations, or co-morbidities increasing complication risk.

Implant procedure results

The implant success rate was 100% (140/140). Type of anaesthesia, anticoagulation, and use of antibiotics were at the discretion of implanters. Sedation and/or local anaesthesia was applied in 93.6% of patients and general anaesthesia used in 6.4%. No systemic anticoagulation was applied in 36%, and various regimens were used in the other patients (40% received heparin and 24% received another anticoagulation method). An anticoagulation antagonist was used to reverse anticoagulation effects in 16 patients (11%). The mean implant time was 37 ± 21 min (range: 11–154 min) with an average fluoroscopy time of 9 ± 7 min. The majority of attempts were successful upon initial device positioning (59%) or two repositioning (22.1%), but the maximum number of attempts in a single patient was 18 deployments (mean ± SD, 2 ± 2). A second device was used in two implants, due to unacceptable electrical measurements. In both cases, electrical measurements with the second device were similar to the measurements of the first device, although ultimately both cases were able to achieve a position with acceptable electrical measurements. While 107 of the 140 devices were placed at the RV apex (76.4%), 33 (24%) were implanted at the anterior septum, mid-septum, or outflow tract (images available in Figure 5). Access site closure was predominantly performed using a suture method (76%), although other methods were observed (such as manual pressure or venotomy occlusion system). The average time to ambulation following the procedure was 13 ± 8 h and the median days from procedure to hospital discharge was 1 day, although day to discharge varied by geography (mean ± SD, 2 ± 2).

Early safety performance

There were no USADEs in the 140 patients, thus the safety objective was met with 100% freedom from USADEs at 3 months (95% confidence interval, 94.0–100%; \( P < 0.0001 \)). Thirty adverse events occurred in 26 patients, all within 17 days of implant (Table 2).

One pericardial effusion was observed in a 90-year-old female who had undergone 18 repositioning because of inappropriate electrical measurements, the highest number of repositioning observed in the study. A pericardial drainage was performed to drain approximately 250 cc of blood, although no tamponade was diagnosed. The same patient experienced an acute myocardial infarction 3 days post-implant and angiography revealed three-vessel coronary artery disease. Transient complete AV block occurred in four patients and resolved within seconds to a few hours. Three cases of transient AV block required pacing via a temporary wire. The fourth resolved with immediate programming to active pacing after device deployment. Each of the four patients experiencing transient AV block had a history of LBBB or prolonged AV conduction (two had second-degree AV block, one had LBBB with first-degree AV block, and one had LBBB). Groin bleeding (‘incision site haemorrhage’) was observed in three of the 140 patients, and a haematoma in two. None of these events were considered serious, and all cases resolved without invasive intervention. There were two cases of arterial pseudoaneurysms. One of the two events was considered serious and required thrombin injection with prolonged hospitalization. The other pseudoaneurysm was not considered serious and resolved without any invasive intervention. There was no apparent relationship to heparin use or closure method approach in the events which were observed at the groin puncture site. (There was a total of 11 events at the groin puncture site and intravenous heparin was administered in approximately half of the events. A suture method was used for closure in each of these 11 cases except one where only manual pressure was applied). One patient death occurred 139 days post-implant, was not cardiovascular related, and was determined to not be related to the procedure or system.
Early efficacy performance

The mean pacing capture threshold at the 3-month visit for the 60 patients measured at 0.24 ms was 0.51 V (95% CI, 0.45–0.56; \( P \leq 0.0001 \)), meeting the efficacy objective. In these 60 patients, the mean electrical values for R-wave sensing amplitude, pacing impedance, and pacing capture threshold at 0.24 ms were, respectively: 11.7 ± 4.5 mV, 719 ± 226 ohm, 0.57 ± 0.31 V at implant, 15.6 ± 4.8 mV, 662 ± 133 ohm, 0.48 ± 0.21 V at 1-month, and 16.1 ± 5.2 mV, 651 ± 130 ohm, 0.51 ± 0.22 V at 3-months (Figure 6). All measurements at all visits were within expected

| Table 1 Patient characteristics |
|-------------------------------|
| **Patient characteristics** | **First 60 (n = 60)** | **Total (n = 140)** | **P-value** |
| Age (years), mean ± standard deviation | 76.8 ± 9.9 | 77.0 ± 10.2 | 0.84 |
| Male, n (%) | 37 (61.7) | 85 (60.7) | 0.86 |
| Pacing indication, n (%) | | | |
| Symptomatic sinus node dysfunction | 20 (33.3) | 40 (28.6) | 0.35 |
| 2nd-degree AV block | 4 (6.7) | 8 (5.7) | 0.72 |
| 3rd-degree AV block | 6 (10.0) | 11 (7.9) | 0.53 |
| SND + AV block | 1 (1.7) | 2 (1.4) | 1.00 |
| AV block with persistent/permanent arrhythmias | 28 (46.7) | 72 (51.4) | 0.39 |
| Other indication | 1 (1.7) | 7 (5.0) | 0.24 |
| Reason for selecting single-chamber pacemaker, n (%) | | | |
| Indications associated with persistent/permanent/chronic atrial tachyarrhythmias | 34 (56.7) | 82 (58.6) | 0.73 |
| Frequent pacing not expected | 21 (35.0) | 45 (32.1) | 0.59 |
| Patients advanced age | 5 (8.3) | 18 (12.9) | 0.21 |
| Significant co-morbidities affecting survival and clinical outcome | 2 (3.3) | 3 (2.1) | 0.58 |
| Previous planned AV nodal ablation | 0 (0.0) | 2 (1.4) | 0.51 |
| Patients expected to be sedentary | 2 (3.3) | 5 (3.6) | 1.00 |
| Patient’s anatomy precludes placement of atrial lead | 1 (1.7) | 2 (1.4) | 1.00 |
| Dual-chamber pacing complication risk deemed to high | 0 (0.0) | 1 (0.7) | 1.00 |
| Other reason | 2 (3.3) | 8 (5.7) | 0.24 |
| Cardiac disease, n (%) | | | |
| Cardiomyopathy, dilated/congestive + Cardiomyopathy, ischaemic | 5 (8.3) | 5 (3.5) | 0.01 |
| Coronary artery disease + Myocardial infarction | 20 (33.3) | 44 (31.4) | 0.72 |
| Pulmonary hypertension | 1 (1.7) | 7 (5.0) | 0.24 |
| Valve dysfunction, tricuspid | 10 (16.7) | 23 (16.4) | 1.00 |
| Hypertension | 46 (76.7) | 111 (79.3) | 0.53 |
| Symptoms, n (%) | | | |
| Congestive heart failure | 5 (8.3) | 13 (9.3) | 0.78 |
| Syncope | 23 (38.3) | 55 (39.3) | 0.86 |
| Comorbidities (%) | | | |
| Diabetes | 17 (28.3) | 34 (24.3) | 0.43 |
| Renal dysfunction | 10 (16.7) | 26 (18.6) | 0.67 |
| Chronic obstructive pulmonary disease (COPD) | 4 (6.7) | 14 (10.0) | 0.39 |
| Implant success rate (%) | 60 (100.0) | 140 (100.0) | 1.00 |
| Procedure duration (min) | | | |
| Mean ± standard deviation | 37.9 ± 24.8 | 36.7 ± 20.7 | 0.54 |
| Fluoroscopy duration (min) | | | |
| Mean ± standard deviation | 10.1 ± 7.8 | 9.1 ± 7.0 | 0.18 |
| Redeployments, n (%) | | | |
| 0 | 34 (56.7) | 82 (58.6) | 0.76 |
| 1–4 | 24 (40.0) | 52 (37.1) | |
| ≥5 | 2 (3.3) | 6 (4.3) | |
| Hours to ambulation following procedure | | | |
| Mean ± standard deviation | 14.7 ± 7.6 | 13.1 ± 8.5 | 0.10 |

\( P \)-value is from the t-test (continuous variables) or Fisher Exact test (categorical variables) comparing those in the early performance cohort (first 60 implanted) to those implanted later.
Paired comparison of the 60 patients from implant to 3-month electrical values demonstrated an increase in R-wave amplitude (4.4 mV, \( P < 0.0001 \)), a decrease in impedance (68 ohms, \( P = 0.006 \)), and a non-significant decrease in pacing capture threshold (0.06 V, \( P = 0.057 \)). Rate response was programmed in 54% (76 out of 140) of patients, programmed to VVIR where an initial testing showed effective rate adaptation to short walking test. Ongoing evaluations in the study will confirm rate response operation via treadmill testing in a subset of subjects.

Ambulatory ECG evaluation

Examination of 24 h ambulatory surface ECG and device electrogram cycle by cycle at the 1-month visit from 25 patients indicated that the device was pacing and sensing as expected. There were no pauses due to inappropriate TPS operation. Additionally, the daily capture threshold testing and hourly threshold confirmation tests were performing as expected.

Longevity estimation

In 60 patients followed to 3 months, cumulative percent pacing ranged from <1 to >99% with a median of 49% (interquartile range, 10.2–75.1%). Based on device use conditions (e.g. heart rate, pulse width, pacing amplitude, impedance) through 3-months, battery longevity was estimated at an average 12.6 years (range 8.6–14.4 years, Figure 7). This estimate does not include

| Apical placement | Mid-septal placement | RVOT placement |
|------------------|----------------------|----------------|
| ![Image of various device positions in RAO view. Left panel: apical device placement; Middle panel: mid-septal device placement; Right panel: right-ventricular outflow tract (RVOT) device placement.](image)

**Table 2** Procedure or system-related adverse events from 140 implanted patients

| 30 total related adverse events | Resulted in death, re-operation, or hospitalization? | Total event rate, n (%) |
|-------------------------------|-----------------------------------------------|------------------------|
| Dysrhythmias                  |                                               |                        |
| Transient atrioventricular block | No                                         | 4 (4, 2.9)             |
| Right bundle branch block     | No                                            | 2 (2, 1.4)             |
| Ventricular tachycardia       | No                                            | 2 (2, 1.4)             |
| Ventricular fibrillation      | No                                            | 1 (0.7)                |
| Events at device placement site | Pericardial effusion without tamponade       | 1 hospitalization prolonged >48 h | 1 (1, 0.7) |
| Acute myocardial infarction   | No                                            | for both events in same patient | 1 (1, 0.7) |
| Pericarditis                  | No                                            | 2 (1, 0.7)             |
| Non-cardiac chest pain        | No                                            | 1 (0.7)                |
| Angina pectoris               | No                                            | 2 (1, 0.7)             |
| Events at groin puncture suite | Arterial pseudoaneurysm                      | 1 hospitalization prolonged >48 h | 2 (2, 1.4) |
| Incision site haemorrhage     | No                                            | 3 (3, 2.1)             |
| Incision site haematoma       | No                                            | 2 (2, 1.4)             |
| Incision site pain            | No                                            | 1 (1, 0.7)             |
| Incisional drainage           | No                                            | 1 (1, 0.7)             |
| Vaso-vagal presyncope         | No                                            | 2 (2, 1.4)             |
| Other                         | Dysuria following procedure                   | No                     | 1 (1, 0.7) |
| Osteoarthritis following procedure | No                    | 1 (1, 0.7)             |
| Back pain during procedure    | No                                            | 1 (1, 0.7)             |
pacemaker-dependent patients and assumes that thresholds remain stable for device lifetime.

**Discussion**

This is the largest data set reported for any transcatheter pacing technology to date. Early results of this pacemaker are encouraging. The TPS demonstrated safety and efficacy through the acute phase by passing both of its early performance assessment objectives. One hundred and forty (100%) successful implants were achieved in a wide variety of patient demographics, cardiac conditions, and indications for pacing. This was accomplished by 37 physicians in 23 study centres in 11 countries. In two procedures, a second device was used due to unacceptable electrical measurements and decision to change the device was motivated by uncertainty due to early experience of implanters. The majority of implants (81%) were successfully completed with one or two positionings.

**Safety performance**

USADE is an ISO standard for assessing primary risks to patient health when implementing a new medical device. The endpoint is often used for safety evaluation, but is uniquely different from the study’s primary objective. The trial had no USADEs after 266.4 months of follow-up, far exceeding its primary safety objective. Thirty adverse events related to the procedure or system were identified in the 140 implanted patients, none of which were due to device dislodgement or infections. There were no deaths related to the procedure or system and no re-operations were required. Telemetry contact remained feasible in all patients. Adverse events did not differ substantially across patient sub-groups.

One case of pericardial effusion occurred in the context of 18 positioning due to undesirable electrical performance. In light of this observation, persistent repositioning should be avoided in order to limit the possible injury to the myocardium and surrounding vessels. In case of undesirable thresholds immediately after deployment, some waiting time should be considered before measuring electricals again. As is frequently observed in tined and helix-based pacemaker leads, thresholds may improve within minutes, thus allowing release of the device. In all cases, consideration of the potential benefit of multiple repositioning vs. risk is warranted.

Transient AV (in patients with LBBB or AV conduction abnormalities) or right bundle branch block during navigation of the delivery tool was reported in a few cases, suggesting a mechanical trauma by the delivery system which bears consideration in crossing the tricuspid valve. Prevention of ventricular arrhythmias warrants a similar consideration. However, these events can occur with any right-heart procedure. Temporary pacing is a preventive option and may be considered in patients with LBBB.

The management of pre- and peri-operative anticoagulants was left to the discretion of the physician because of lack of experience with the implantation of the TPS. No specific recommendation was
given to investigators who were free to define their own protocol with the condition that all catheters were flushed with heparinized saline, and that a heparinized drip was placed on the introducer to reduce clotting. A variety of approaches were used and minimal haematoma or bleeding were observed, without correlation to approach (40% received heparin and 36% did not receive any anticoagulation; there were two haematomas and three events of minor groin bleeding). Therefore, no obvious advantages or disadvantages due to anticoagulation approach have become apparent, although a heparin bolus may reduce risk of clotting on the device and within the delivery tool. The risk of inadvertent arterial puncture (with possible subsequent complications such as pseudoaneurysms, AV fistulas, and haematomas) might be mitigated by using ultrasound techniques for venous access, although this has not been evaluated within this study. With regard to post-procedural closure/haemostasis, investigators have successfully used various techniques as a pre-specified closure method was not mandated. The majority (46%) of closure methods utilized manual pressure with a suture method (Figure of 8, purse string). A suture method without manual pressure was used in 39% of cases. A vascular closure device was used in 11% of cases (with or without pressure or a suture), and manual pressure alone was used in 4% of cases.

Efficacy performance

Device functionality and efficacy was successfully demonstrated at 3 months with a mean pacing threshold of 0.51 V at 0.24 ms pulse width; exceeding the intended goal of 2.0 V. Further, ambulatory ECG monitoring in the first 25 patients at 1-month demonstrated the device was performing as expected. Pace thresholds remained low and stable, with no measurements at or above 2.0 V at 3 months follow-up. This last observation is important because any programming requiring high energy levels may result in a significant reduction in the life expectancy of the device. One patient had a pacing threshold >2 V at implantation, though this decreased to <1 V at 3 months. Based on pacing conditions from the 60 patients followed to 3 months, mean device longevity was estimated at 12.6 years, with 95% of the patients over 10 years and shortest of 8.6 years. This longevity performance would be no worse than conventional pacing systems.14,15 The TPS uses a new capture management approach with automatic hourly 0.5 V safety margin confirmations to ensure pacing outputs remain at safe levels and to optimize battery longevity.

Alternate transcatheter pacemakers

While other manufacturers are developing transcatheter pacemakers, it will be of importance to consider technical differences in the design and performance of the various technologies. In the LEADLESS Trial, the Nanostim™ leadless cardiac pacemaker (St Jude Medical, St Paul, MN, USA) was successfully implanted in 32 of 33 patients.16 One patient experienced a RV perforation and cardiac tamponade during the procedure, and later died as a result of stroke. Two patients required device retrieval post-implantation, one for inadvertent left-ventricular placement and the other for developing an indication for an implantable cardioverter defibrillator. Compared with TPS, the Nanostim device is of similar size but longer (41.4 vs. 25.9 mm) and narrower in diameter (18 vs. 20 Fr). Rate response is controlled by RV blood temperature compared with an accelerometer in TPS. Although the implant approach is also transcatheter delivered via the femoral vein, endocardium fixation is different. The TPS uses four protractible nitinol tines vs. a fixed helical coil with Nanostim.

Technology implications

Transvenous lead technology was developed in 1959 and with the introduction of implantable pacemakers in the 1960s, the implant technique has largely been unchanged. Today, it is estimated that ~600 000 pacemakers are implanted worldwide each year,17 358 000 of which are in the USA.18 Compared with single-chamber atrial-based systems and the more costly dual-chamber systems, VVI pacemaker utilization varies regionally with rates reported from trials of 8% in the USA19 and 25% in the Netherlands.20 VVI pacemakers are guideline recommended for patients with permanent or persistent atrial arrhythmias and slow intrinsic heart rates.20 Future developments of this technology may include AAI, VDD, DDD, or even biventricular pacing systems. However, in the meantime, mode selection may be influenced by the technological advantages of a miniaturized transcatheter pacemaker, advantages that include the absence of lead and pocket related risks and device implant cosmesis. Atrial and dual-chamber pacing modes have recently been preferred for bradycardia patients without permanent or persistent atrial tachyarrhythmia due to risks associated with AV dysynchrony, notably pacemaker syndrome.21 Over one-third of the patients who received TPS in our trial were without history of permanent or persistent atrial tachyarrhythmia. Of these patients, the primary pacing indication was SND (45%) or AV block (39%). Physicians selected VVI pacing for reasons due to ‘infrequent pacing expected’ (69%), ‘advanced age’ (22%), and other reasons that included sedentary lifestyle, anatomical limitations, or co-morbidities increasing risk of complication.

Limitations

This study is non-randomized and so the evaluation of benefit of this new technique over contemporary single-chamber or dual-chamber systems is indirect. This is the first report of early performance of the TPS in a multi-stage assessment protocol and there was limited representation of pacemaker-dependent patients in this cohort. The patient cohort is too small to determine any recommendation regarding anticoagulation management and post-procedural closure/haemostasis. No devices have yet to be retrieved, thus retrievability remains uncertain. Also, performance with multiple or concomitant devices has not yet been observed. This is an ongoing study that may bring more precision regarding these issues.

Conclusion

Early performance assessment shows the TPS pacemaker can safely and effectively be applied. It is premature to draw definitive conclusions about the benefits of this system. Long-term safety and benefit associated with the absence of a subcutaneous pulse generator and transvenous lead will further be evaluated in the trial. However, these early results meet and exceed initial pre-specified expectations. The fact that such data were obtained in a multicentre study involving 37 investigators at 23 different centres around the world with a variety of patients is encouraging.
Supplementary material

Supplementary material is available at European Heart Journal online.

Acknowledgements

We would like to thank the physicians who contributed to the early performance cohort as listed in the Supplementary material online. We would like to thank Eric Williams for his technical support and Harrison Hudnall for his editorial support in the preparation of this article.

Funding

This work was supported by Medtronic plc as the Sponsor of the Micra Transcatheter Pacing Study. Funding to pay the Open Access publication charges for this article was provided by Medtronic plc.

Conflict of interest: P.R.: consulting fees/ honoraria; G.Z.D.: consulting fees/ honoraria; C.S.: consulting fees/ honoraria; K.S.: consulting fees/ honoraria; L.C.: consulting fees/ honoraria; S.Z.: consulting fees/ honoraria; L.V.A.B.: consulting fees/ honoraria; R.E.K.: consulting fees/ honoraria; R.O.: consulting fees/ honoraria, speaker’s bureau; L.M.: consulting fees/ honoraria; C.S.: consulting fees/ honoraria; K.S.: consulting fees/ honoraria.

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