Initial experience at one Swiss centre

Direct His bundle pacing in routine clinical practice

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Summary

Direct His bundle pacing has recently attracted interest as a more physiological alternative to right ventricular or biventricular stimulation. The advent of new tools has facilitated the implantation procedure. This report relates our initial experience with this technique in our first 50 patients.

Key words: direct His bundle pacing; pacemaker; implantable cardioverter defibrillator; cardiac resynchronisation therapy

Introduction

It is well established that chronic right ventricular (RV) apical pacing may have an adverse effect on left ventricular systolic function [1, 2], leading in the long term to adverse clinical outcomes such as heart failure [3], atrial fibrillation [4, 5] and even death [6]. In order to avoid these adverse effects, the interventricular septum or right ventricular outflow tract have been proposed as alternatives to RV apical pacing, but results have been equivocal [7, 8]. In the quest for more physiological alternatives, His bundle pacing (HBP) was first performed by Desmukh et al. in 2000 [9]. This technique has the advantage of avoiding the electrical (and thereby mechanical) dyssynchrony induced by myocardial pacing by recruiting the intrinsic conduction tissue to activate the ventricles, resulting in a narrow QRS complex. Furthermore, it was shown by Narula in 1977 that pacing of the His bundle can correct bundle branch block (BBB), implying a proximal site of conduction disturbance with longitudinal dissociation within the His bundle [10]. HBP may therefore be used in lieu of cardiac resynchronisation therapy (CRT), for example in patients with failed coronary sinus lead implantation.

Despite its virtues, the technique was not readily adopted because of the complexity of the procedure, which required mapping of the His bundle using a diagnostic electrophysiology catheter, and positioning of a standard pacing lead with a manually curved stylet. Procedures lasted hours and resulted in high capture thresholds. The advent of a steerable sheath to deliver a 4.1 F lumenless lead, the Select Secure 3830 model (Medtronic, MN) facilitated the procedure (fig. 1), but capture thresholds remained high (on average $2.3 \pm 1.0$ V/0.5 ms) [11]. We have been using this lead for pacemaker implantation in paediatric patients for over a decade [12]. More recently, a sheath with a fixed curve specifically designed for locating the His bundle (Medtronic Select Site C315 catheter – see figure 1) has further facilitated the procedure by obviating the need for a diagnostic catheter, and with improved thresholds (mean $1.35 \pm 0.9$ V/0.5 ms) [13].

In a nonrandomised comparison of 304 patients with HBP and 433 patient with right ventricular pacing, the former group had significantly fewer heart failure hospitalisations and a trend towards reduced mortality [14]. Two small randomised cross-over studies (combining a total of 50 patients) showed that HBP was safe, with a tendency to better left ventricular ejection fraction (LVEF) compared with right ventricular pacing [15, 16]. In a randomised cross-over study in 21 patients (12 with complete data) HBP performed similarly in terms of LVEF, New York Heart Association (NYHA) class, quality of life and 6-minute walk distance compared with biventricular pacing [17].

Following these encouraging developments, and magnetic resonance imaging (MRI)-conditional labelling of the 3830 lead in May 2017, the first patients were implanted with HBP at our institution. This article reports our experience in our first 50 patients.

Material and methods

Patient population

Consecutive patients with a standard indication for pacing according to current guidelines [18], in whom frequent ventricular pacing was anticipated and in whom HBP was attempted by a single operator (H.B.), were included. Patients were recruited from the cardiology department of the University Hospital of Geneva. The protocol was approved by the institutional ethics committee, and patients gave informed consent to participate in the study.
Implantation technique

HBP was performed according to implantation techniques previously described [19, 20], using the Medtronic 3830 lead with the C315 His delivery catheter (or optionally, if unsuccessful, the C304 deflectable catheter). The His lead was implanted first (this allowed the 3830 lead to be used for RV or right atrial pacing instead, if HBP was unsuccessful), unless the patient was scheduled for CRT implantation (in this case, the coronary sinus lead was implanted first). In all cases, the His bundle was mapped in the right anterior oblique (RAO) 30° view using the pacing lead, in a unipolar sensing configuration (as the ring electrode was usually covered by the guiding catheter during mapping). The lead was connected to the atrial channel of the pulse sense analyser (PSA) at 50 mm/s sweep speed and maximum amplification (0.05 mV/mm). With experience, it was noticed that His potentials were more readily mapped on the atrial aspect of the annulus, and we avoided crossing the tricuspid valve (observed as a “jump” of the catheter). If no His potentials were visible on the programmer screen, a printout was obtained as this occasionally allowed visualisation. The Medtronic PSA was replaced by the Abbot programmer (using unfiltered signals) for the last 10 cases, as the display is larger (fig. 2), and having two PSAs facilitated backup pacing with continued mapping in the event of atrioventricular block. The Medtronic PSA was, however, always used for measuring sensing thresholds using the “new” filter settings, which reflect those of the corresponding channel of the implanted generator. If no His potentials were visible, pace-mapping was performed. The lead was fixated by twisting the body until a slight buildup of resistance was felt. To improve tactile feedback, a transvalvular introducer tool was inserted in the guiding catheter valve. A standard 12-lead ECG was recorded during lead testing in all instances to confirm His capture, starting at 10 V/0.5 ms output and at 1 V decrements to evaluate transitions in QRS morphology, for example from nonselective HBP (with a “pseudo-delta” wave corresponding to local myocardial capture, see fig. 3), to selective HBP (with an isoelectric interval in all 12 leads between the pacing spike and QRS onset) as well as correction of BBB (fig. 4). In the event of HBP without correction of BBB, the lead was repositioned in a slightly more distal (cranial and more ventricular) position. “Para-Hissian” pacing without evidence of capture of conduction tissue (i.e., no transition in QRS morphology with decreasing output) and selective HBP without correction of LBBB, were considered failures. Backup ventricular pacing was provided in all cases. In patients with chronic atrial fibrillation (AF), the His bundle lead was connected to the atrial port of the pacemaker / implantable cardioverter defibrillator (ICD). The device was programmed to a DDDR mode with a paced atrioventricular delay of 140 ms and inactivation of ventricular safety pacing after checking for crosstalk. In patients in sinus rhythm, the His bundle electrode was connected to the left ventricular (LV) port of a CRT pacemaker/ICD (fig. 5). The device was programmed to a DDD mode with sequential biventricular pacing and the maximum delay of 80 ms (left ventricle first).

A maximum of 30 minutes was allocated for mapping the His. If no His potential was found, or pace-mapping was unsuccessful, His lead implantation was abandoned, as was also the case if thresholds were unacceptably high (based upon clinical judgment for each...
patient individually). In these instances, the 3830 lead was implanted in the right ventricle or the right atrium, as required.
Details on His lead positioning (e.g., material used, numbers of attempts, visualisation of the His potential and a current of injury, thresholds, etc.) were recorded. Electrical parameters at routine follow-up were also captured.

**Statistical analysis**
The IBM SPSS v24 program (Armonk, NY, USA) was used for calculations. Data are expressed as mean ± standard deviation unless specified otherwise. Fisher’s exact test was used for comparing proportions. A p-value <0.05 was considered significant.

*Figure 3:* His bundle pacing in a patient in atrial fibrillation with normal baseline QRS. The tracing shows nonselective His capture, alternating with selective His capture with disappearance of the pseudo-delta wave (shown by the asterisks in lead aVL).

*Figure 4:* Electrocardiogram in a patient with baseline left bundle branch block. Transition (shown by the asterisk) between nonselective His bundle pacing with correction (first four beats) followed by selective His bundle capture without correction and widening of the QRS complex corresponding to left bundle branch block.
**Results**

A total of 50 patients were recruited over a 1-year period (between 8 May 2017 and 4 June 2018). Patient demographics are shown in Table 1. The youngest patient was 32 years old and had congenital atrioventricular (AV) block, and the oldest was aged 91 years and required device implantation for an “ablate and pace” strategy for rapidly-conducted AF with heart failure and systolic dysfunction.

**Procedural success**

Overall, 33/50 (66%) had successful HBP implantation (Fig. 6). All procedures were performed using left-sided access, apart from four cases, of which one failed (handling of the C315 guiding catheter from the right side was not difficult). A His potential was recorded by the pacing lead in 39/50 (78%) of cases, and pace-mapping led to successful location of the His bundle in 5/11 (45%) of the remaining patients. HBP was successful in 21/25 (84%) of patients with a normal baseline QRS, compared with 12/24 (50%) of patients with BBB ($p = 0.016$). A patient in AF with paced ventricular rhythm had failed HBP due to unsuccessful localisation of the His bundle – all six other patients who were in complete AV block at the time of the procedure had successful HBP implantation. The C304 deflectable catheter was used in five patients and led to successful implantation of the lead in one of these patients.

The average time taken to locate the His (in cases where this was possible) was $6.3 \pm 5.9$ minutes, and the total implantation duration attributed to the His lead (in all

| HBP success | n = 33 (66%) |
| HBP failure | n = 17 (34%) |
| Selective HBP | n = 15 | | Non-selective HBP | n = 17 |
| No His / pacemapping unsuccessful | n = 6 | | Selective HBP without correction of LBBB | n = 5 |
| Fixation unsuccessful | n = 3 | | High threshold | n = 3 |

**Figure 5:** Chest X-ray in a patient implanted with a pacemaker for atrioventricular block. RA: right atrial lead; His: His bundle lead (connected to the left ventricular port of the biventricular pacemaker); RV: right ventricular lead on the interventricular septum (connected to the right ventricular port of the pacemaker).

**Figure 6:** Summary of procedural success of His bundle pacing. HBP: His bundle pacing, LBBB: left bundle branch block.
procedures – also those with failed implantation, and including the time required for lead testing, which is considerably longer than for standard leads owing to ECG analysis) was 26.6 ± 17.0 minutes.

The His lead was connected to the atrial port in 20 patients (all in AF/flutter) and to the LV port in the remaining 13 patients (all in sinus rhythm). Of the 33 patients with successful HBP implantation, a pacemaker was implanted in 26 patients and an ICD in seven subjects. The only complications noted were transient complete AV block in three patients who had left bundle branch block (LBBB), and new right bundle branch block (RBBB) after the His lead was fixed in one patient with a normal baseline QRS.

Electrocardiographic parameters
Overall, capture of the His was possible in 40 (80%) patients, of whom 21 showed selective capture and 19 nonselective capture. Of the 23 patients with BBB in whom His capture was possible, 10 (43%) had correction of the QRS duration by at least 20 ms. Of these patients, 5/11 had RBBB and 5/11 patients had LBBB (p = 1.0 for the comparison). The patient with nonspecific intraventricular conduction delay did not have correction of the BBB. Of note, three of the four patients with Medtronic CoreValves had LBBB (one had normal QRS morphology), and none had correction of intraventricular conduction delay by HBP (only selective His capture without correction). Localisation of the His bundle was very easy (within 5 minutes) in all four patients and landmarked by the inferior corner of the valve cage in the RAO view.

Overall, the paced QRS duration in the 33 patients with successful HBP was 108 ± 18 ms.

Electrical parameters
His bundle current of injury after lead fixation, associated with reduced capture thresholds [21], was observed in 11/39 (28%) of patients with a visible His potential. The capture threshold of the His bundle with unipolar pacing was 1.7 ± 1.1 V/0.5 ms (range 0.3–4.6 V/0.5 ms). Bipolar sensing was 4.2 ± 4.4 mV (range 0.5–8.7 mV). Unipolar impedance was 567 ± 173 Ohms. Follow-up was available for 25 patients after a duration of 3.1 ± 2.8 months. Pacing polarity was programmed to unipolar or extended bipolar (except if the His lead was connected to the atrial port of CRT-Ds, as only bipolar pacing is available in these devices). In patients with the His lead connected to the atrial port, sensitivity was programmed to 4 mV as ventricular sensing was ensured by the backup right ventricular lead (and to avoid oversensing of atrial or His potentials). HBP capture thresholds were 1.2 V ± 1.1 V at a pulse width of 0.5 ± 0.2 ms; sensing was 3.2 ± 3.3 mV and impedance was 385 ± 110 Ohms.

Discussion
The main findings of this report are that (1) HBP is successful in two thirds of patients with a higher success rate in patients with normal baseline QRS (84%) than in those with BBB (50%), (2) Correction of QRS by >20 ms in patients with BBB is possible in almost half of the patients and (3) Electrical parameters are acceptable and stable over short-term follow-up. Reported success with HBP in experienced hands was initially reported as 80% using current tools [13] and is now >90%, which is considerably higher than the 66% in this report. Reasons for this are that (1) this is our initial experience, including the learning curve, (2) a maximum of 30 minutes was allowed for positioning the His
lead and success rates are likely to have been higher with more allocated time, (3) strict criteria were used for defining success (e.g., “para-Hissian” pacing, included in some reports, was considered as failed implantation), (4) the setup in the operating room was not optimal (e.g., an electrophysiology bay and large-screen display were not available, and would have greatly facilitated mapping of His potentials), (5) no diagnostic catheters were used for locating the His bundle, contrary to some practice [23], and (6) the patient population was relatively complex (a high proportion with conduction disorders and comorbidities, which may have affected success rates).

It is interesting to note that patients with BBB had a lower success rate than those with a normal baseline QRS. One reason for failed implantation was His capture without correction of LBBB. The proportion of patients in whom this can be achieved is as yet not well defined. In 1977, Narula et al. [10] reported for the first time correction of LBBB by distal His pacing, with correction in 25/80 (31%) patients, but only temporary pacing with diagnostic catheters (as opposed to screw-in leads) were used. Ajijola et al. [23] studied 21 patients who received HBP in lieu of biventricular pacing, of whom 15 (71%) had QRS narrowing by >20%. In our series, the proportion of patients with BBB correction by >20 ms was only 10/23 (43%), but might have been higher with more experience. The three patients with transcatheter aortic valve implantation (TAVI) and LBBB who all had failed implantation due to selective HBP without correction are an interesting group because conduction disease may be more distal in this setting than with degenerative disease. A recent publication on HBP in 30 patients with prosthetic valves reported a success rate of 93% [24]. This series included four patients with TAVI (all Edwards Sapien valves), in two of whom HBP was successful. Our series reports for the first time HBP in patients with Corevalves, which extend lower and may result in a more distal block, explaining the failure to correct LBBB, but this warrants further study.

Capture thresholds were acceptable and comparable to previous reports [25, 26], and remained stable at short-term follow-up. We chose to implant a backup ventricular lead in all patients, as this was our initial experience, and also because increase in capture threshold and requirement for lead revision has been reported in 6.6% of patients [25]. Furthermore, sensing amplitudes were sometimes as low as 0.5 mV, and the backup ventricular lead also served to avoid sensing issues. However, this comes at an extra cost of an additional lead, and of a dual-chamber instead of a single-chamber generator or a biventricular instead of a dual-chamber device. Since our first 50 patients, we have started implanting HBP without backup ventricular pacing in cases with good capture and sensing thresholds, and if atrioventricular nodal ablation is not planned. It should be noted that in the case of CRT implantation in patients in chronic AF, the atrial port is usually plugged, and HBP is an option that comes with the marginal extra cost of an additional lead.

The recent surge in interest in HBP is spurring the development of new tools in this field. A deflectable delivery catheter with a posterior angled tip will facilitate mapping of the His bundle and should be available in the near future (in the meantime, we have started using the deflectable catheter for coronary sinus access, coupled with a foreshortened Attain Select II 90° inner catheter). Delivery catheters to accommodate standard 6 F stylet-driven leads with extendible screws are also being developed by competitors. Hopefully, further developments in lead design (e.g., with a longer helix) will further facilitate implantation and lower capture thresholds.

Study limitations: The number of patients and follow-up duration are relatively limited, and the full scope of issues encountered with HBP is not covered by this report.

Conclusions

HBP has been performed at our centre for over a year with good results and has now become routine clinical practise in patients requiring frequent ventricular pacing or in the case of CRT and chronic AF (in whom a His lead is connected to the atrial port). The technique has also been adopted this year by several other Swiss centres. Much as with CRT over a decade ago, the advent of new tools has facilitated the procedure and allowed it to enter mainstream clinical practise. The tools will no doubt continue to evolve and further facilitate implantation, and future research will determine which patients benefit the most.

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Disclosure statement

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