ACUTE HIS BUNDLE INJURY DURING PERMANENT HIS BUNDLE PACING: MECHANISTIC INSIGHTS INTO INTRA-HISIAN BLOCK

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ABSTRACT. Permanent His bundle pacing (HBP) is a physiological alternative to right ventricular pacing. Acute injury to the His bundle can occur during permanent HBP in the form of transient or permanent bundle branch blocks. Observations of mechanical injury to the His bundle during HBP lead fixation and electrophysiological response to HBP provides additional mechanistic insights into the pathology of bundle branch block and intra-Hisian atrioventricular block.

KEYWORDS. Bundle branch block, HV block, intra-Hisian block, left branch block, permanent His bundle pacing, right bundle branch block.

Introduction

Permanent His bundle pacing (HBP) is a physiological alternative to right ventricular (RV) pacing. RV pacing is associated with ventricular dyssynchrony, left ventricular systolic dysfunction, heart failure, and increased mortality.1-5 Despite the technical challenges during early experience with permanent HBP,6 it is now clear that this form of pacing is feasible in routine clinical practice.7,8 While permanent HBP is intuitively successful in patients with atrioventricular (AV) nodal block and preserved His–Purkinje conduction, recent studies report the feasibility of HBP to correct the conduction disturbances in patients with infranodal AV block and in patients with bundle branch blocks (BBBs).9,10 Early studies in animals and humans have shown that BBBS and infranodal AV blocks are due to lesions within the main His bundle.11-13 This article will review the incidence of acute injury to the main His bundle during permanent HBP and how it helps to understand the mechanisms of intra-Hisian conduction disease.

His bundle injury current

Permanent HBP utilizes the placement of an active fixation pacing lead at the His bundle region. Early in HBP experience, a traditional stylet-driven lead with retractable or exposed screw was used.5,14 HBP thresholds obtained using these leads were significantly higher (>2 V) than RV capture thresholds. In addition, the fluoroscopy and procedural duration for HBP were significantly longer than RV pacing because of the inability to precisely position the electrode in the His bundle region. With the availability of a dedicated pacing lead (SelectSecure 3830™, Medtronic Inc., Minneapolis, MN) and delivery sheaths, the HBP electrode can be accurately positioned in or near the His bundle region. This pacing lead is 4.1 Fr in diameter with no inner lumen and has an exposed screw of length 1.8 mm. The distal electrode screw is steroid eluting to allow for improved pacing thresholds. A recent study demonstrated that an acute His bundle injury current can be recorded from the unipolar or bipolar HBP lead electrograms at the time of implant in about 40% of the patients undergoing HBP (a series of 60 consecutive patients).15 In these patients the His bundle capture thresholds were significantly

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lower than in the patients in whom His bundle injury current was not recorded. With the currently available lead and delivery systems, His bundle capture thresholds of <1.5 V can consistently be achieved in the majority of patients. Recording of acute His bundle injury current during implant confirms that the lead is located directly on the His bundle and the electrode screw has penetrated the outer fibrous insulation of the His bundle itself, enabling better pacing thresholds. On the other hand, in the remaining 60% of the patients where a His bundle injury current was not recorded, it is likely that the electrode was in the vicinity of the His bundle without penetrating its fibrous sheath. It is also likely that in some patients the His bundle is deeper and could not be reached because of the short electrode screw length. Enhanced mapping techniques and leads with longer screws may help achieve more precise positioning of the HBP lead to achieve HBP thresholds similar to myocardial capture thresholds.

Injury to the His bundle

Catheter manipulation during electrophysiology study or cardiac catheterization is known to be associated with transient or permanent BBB. Targeted positioning of a permanent active fixation pacing lead in the His bundle region is expected to cause injury to the His bundle during implantation. A recent study demonstrated that during permanent HBP in 358 patients without pre-existing His–Purkinje conduction disease, acute injury to the His bundle in the form of bundle branch blocks or AV block developed in 7.8% of the patients. RBBB developed in 5.8%, LBBB in 0.8% and complete His-ventricular (HV) block occurred in 1.1% of the patients (Figures 1 and 2). The majority of these conduction disturbances occurred during active fixation of the lead, suggesting acute injury to the His bundle. In addition, when these conduction disturbances occurred during lead fixation, an injury current was invariably noted in the His bundle electrogram, confirming penetration of the electrode beyond the fibrous insulation into the His bundle itself. In 2.5% of patients, the RBBB persisted and did not resolve during follow-up. However, in the remainder of patients, the injury to the His bundle was transient with complete recovery of BBB and HV block. Development of BBB or complete AV block during permanent HBP has several clinical implications. If permanent, His bundle pacing is attempted in patients with LBBB transient or permanent RBBB could lead to complete AV block. Similarly, complete AV block can also occur in patients with normal His–Purkinje conduction. It is essential to be prepared for temporary transcutaneous or transvenous pacing. If the AV block occurred during lead fixation, pacing from the HBP lead would invariably capture the distal His bundle. Alternately, additional venous access and pacing electrode (atrial) should be readily available for transvenous pacing to avoid catastrophic consequences from prolonged asystole. Careful manipulation of the sheath with gentle movements is essential to prevent injury to the His bundle during mapping. It is preferable that mapping for the His bundle is performed from the atrial to the ventricular side of the tricuspid annulus to avoid injury to the distal His bundle early during mapping.
In addition, if the proximal His bundle is injured, it is still possible to fix the lead in the distal His bundle.

**Mechanistic insights**

Acute current of injury demonstrated as ST-segment elevation during myocardial lead placement is due to focally damaged cell membranes consequent to the trauma of electrode pressure against the myocardium. The resulting edema leads to an acutely higher pacing threshold of the tissue in contact but invariably improves over the next several minutes. The current of injury is a marker of excellent tissue contact and hence results in better pacing thresholds. This holds true in the setting of permanent HBP. In patients with His bundle injury current, the acute His bundle capture threshold may be high (>2 V) initially but then improves frequently to <1 V by the end of the procedure because of resolution of the tissue edema.

Acute BBB and complete HV block that occur during HBP lead implant are most likely due to severe edema resulting from direct trauma during the lead fixation. As previously described, a His bundle injury current is recorded on the HBP lead electrograms in all these cases. The development of RBBB or LBBB during the lead placement in the main His bundle, correction of the induced BBB during pacing at the site, and capture of the entire His bundle during pacing confirms that discrete lesions in the main His bundle can lead to right or LBBB. Early investigators in dog models have previously demonstrated similar observations. Apart from inducing His bundle injury (intra-Hisian block) by ligating the first septal artery and causing myocardial ischemia in canine models, El-Sherif et al. were able to induce localized damage with a needle point inserted into the His bundle while action potentials were being recorded from microelectrodes proximal and distal to the damaged zone causing BBBs. These findings reversed after several minutes.

During permanent HBP attempts in more than 600 patients in our laboratory, we have observed six cases of complete AV block. In four of these patients there was evidence for complete HV block (Figure 3), injury current on the His bundle electrograms and idioventricular, wide-complex escape rhythms. In these patients, conduction normalized within 10 min to 24 h, and His bundle capture thresholds were less than 1 V. In two of these patients there was evidence for split His potentials and His bundle escape rhythms with narrow QRS (Figure 4). His bundle capture thresholds were acutely high (>3 V) and improved to less than 0.5 V within 24 hours. In one patient, conduction did not recover at 24 hours but had completely normalized at the 2-week follow-up. In a recent report of permanent HBP in patients with advanced AV block, in 29 of 54 (54%) patients with infranodal disease, the His bundle could be successfully recruited by pacing at the site with evidence for HV block on a local electrogram recorded from the pacing lead. The intracardiac electrograms from the HBP lead in these patients is similar to those observed during acute

Figure 2: (a) Baseline electocardiogram with intracardiac electrogram from the His bundle pacing (HBP) lead before lead fixation. H: His. (b) Following fixation of the HBP lead, there is a significant current of injury (arrow) on the His bundle electrogram with new left bundle branch block (LBBB). (c) Pacing from the HBP lead at decreasing output (from 1.5 to 1 V). The first two beats show non-selective His bundle capture with correction of LBBB. The third beat (*) shows selective left bundle capture with right bundle branch block morphology. The next three beats show non-selective left bundle capture with right ventricular fusion. LBBB resolved within 30 minutes.
HV block induced during lead fixation (Figure 5). The ability to correct HV block in both these scenarios suggests that the intra-Hisian lesion causing HV block is fairly discrete, and virtual electrode polarization created during pacing is able to capture the His beyond the site of block.22,23 In the series of HBP in infranodal AV block,
we reported two patients with split His potentials. A similar clinical scenario was noted during the acute HV block induced during HBP in two patients where split His potentials developed following lead fixation (Figure 4). The lesions causing intra-Hisian block with split His potentials are very discrete and pacing at these sites can overcome the conduction block. In the above-mentioned series of infranodal AV block, in 12 of 54 (22%) patients pacing at sites demonstrating HV block did not result in distal His–Purkinje conduction. Advancing the lead slightly more distally resulted in His capture and conduction. It is likely that in these patients, the intra-Hisian disease was extensive or the initial lead placement was more proximal and did not result in His capture. During HBP lead implantation, we have also observed five cases of transient LBBB that resolved over 5 min to 24 h. Pacing from the HBP lead corrected the LBBB at low outputs. Narula et al.11 had demonstrated that LBBBs are often due to longitudinal dissociation in the main His bundle with resultant asynchronous conduction delay. In addition, pacing the distal His region could correct these conduction disturbances. LBBB during HBP lead placement at the right AV septal region confirms these early observations. Correction of the LBBB by pacing at the site causing LBBB also attests to the discrete nature of conduction disease in the main His bundle leading to LBBB.

In the majority of patients with acute injury to the His bundle, conduction disturbances resolve. However, RBBB persisted in 2.5% of patients during follow-up. While it is conceivable that acute injury may lead to future conduction disturbances in these patients, we have not observed new BBB or AV blocks during medium-term follow-up.

Conclusions
Injury to the His bundle may occur during permanent HBP. Careful manipulation of the sheath and the lead with gentle movements is essential to prevent His bundle injury during implantation. These conduction disturbances in the form of BBB or complete HV block provide mechanistic insights into the pathophysiology of intra-Hisian conduction disease and offer partial explanation as to how permanent HBP can correct these conduction abnormalities. This phenomenon of His bundle injury has to be taken into consideration while designing newer leads with longer screws to improve HBP outcomes.

References
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