His bundle recording and electrical stimulation of atria in patients with Wolff-Parkinson-White syndrome type A

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Two patients presenting with WPW syndrome type A were studied by His bundle potential recording, endocavity electrocardiogram, and electrical stimulation of right and left cardiac cavities.

The beginning of the ventriculogram preceded the appearance of the His bundle potential confirming the pre-excitation of the ventricles by another conduction system than the normal one.

Increasing the prematurity of coupled right atrial extrasystoles increased the degree of delay on the normal pathway and therefore increased the degree of pre-excitation. Both pathways contributed to ventricular depolarization, as shown by an increasingly longer total depolarization time.

The pre-excitation was more obvious with left atrial stimulation. Ventricular depolarization seemed to be completely taken in charge by the anomalous pathway allowing no time for excitation from normal pathway as shown by a constant total depolarization time. In one patient, the atrio-bundle of His conduction was shorter with left atrial than right atrial stimulation.

During bouts of tachycardia, which were more easily induced from the left cavities than from the right, retrograde conduction was faster to the left than to the right atrium.

Our findings are consistent with an anomalous pathway lying preferentially on the left side of the heart in WPW syndrome type A.

The present conception of the Wolff-Parkinson-White (WPW) syndrome depends on the existence of accessory atrioventricular (AV) fibres short-circuiting the normal nodal pathway. This concept has been reinforced by new methods of cardiac exploration developed during recent years. Thus the recording of the potential of the His bundle by cardiac catheterization has confirmed the reality of the short circuit of the His bundle in WPW syndrome (Castellanos et al., 1970; Castillo and Castellanos, 1970; Coumel et al., 1970; Touboul et al., 1971). Electrical stimulation of the heart has also enabled a better understanding of the mechanism of the pre-excitation (Lau et al., 1967; Durrer et al., 1967; Puech et al., 1968). Recently Wellens, Schuilenberg, and Durrer (1971) have reported the results of an investigation of WPW syndrome type A by left atrial stimulation. These authors have found strong arguments in favour of a localization of the accessory pathway on the left side of the heart.

We had the opportunity of performing a study of the pre-excitation in 2 patients presenting a WPW syndrome type A using the His bundle recording and the stimulation of the left atrium and the right atrium.

Patients and methods
In the Table are presented the sex, ages, types of WPW syndrome, and modalities of the tachycardia of the two patients studied. Electrocardiograms are presented in Fig. 1 and 2. They fulfill the criteria of WPW syndrome type A in Case 1

<table>
<thead>
<tr>
<th>Case No.</th>
<th>Sex</th>
<th>Age (yr)</th>
<th>WPW (type)</th>
<th>Modalities of tachycardia</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>F</td>
<td>37</td>
<td>A</td>
<td>Paroxysmal atrial fibrillation and regular tachycardia intermittently for a week</td>
</tr>
<tr>
<td>2</td>
<td>F</td>
<td>60</td>
<td>A with LAD</td>
<td>'Palpitations' for six years</td>
</tr>
</tbody>
</table>

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FIG. 1 Case 1. WPW syndrome type A.

FIG. 2 Case 2. WPW syndrome type A with left axis deviation.

FIG. 3 Case 1. Coupled right atrial stimulation. To read the ordinate, add 100 for SH− SR− and 200 for SJ−. See terminology for the definitions of the different intervals. Shortening the PS increases the SH− interval and in parallel the SJ−. SR− is constant. With very short PS (250 msec) there is an intra-atrial delay (longer SP−) responsible for the lengthening of the other intervals.
and type A with left axis deviation (LAD) in Case 2, according to Rosenbaum, Hecht, and Wilson (1945).

Patients were premedicated with pentobarbitone 120 mg in suppository form. His bundle recording was performed according to the methods of Scherlag et al. (1969). Under local anaesthesia, a bipolar catheter1 (distance between the electrodes: 10 mm) was introduced by the Sel-dinger technique through the right femoral vein and fluoroscopically positioned across the septal leaflet of the tricuspid valve where His bundle activity (H) could be recorded as a biphasic or triphasic deflection of 15 to 20 msec duration. Using the same technique in the left femoral vein, a multipolar catheter2 (5 pairs of electrodes: 2 mm distance between the electrodes and 10 mm between the pairs) was positioned in the right atrium. Later it was manoeuvred across an open foramen ovale in the left atrium. The tip electrodes were used to deliver the electrical stimuli and the next pair of electrodes to record the potentials. A careful sketch of the positions of the electrodes was done: in the right atrium the tip electrodes were in contact with the sinus node region and the other electrodes near the external border of the right atrium. In the left atrium the tip lay near the atrial appendage and the proximal electrodes curved in the superior third of the atrium.

The different leads were connected to a 4-channel direct ink recorder3 via an incorporated distribution switch box: filtering and preamplification were applied to the His lead derivation. Paper speeds were 50 or 100 mm/sec. Rapid atrial pacing was done using a Medtronic model 5841 pacemaker. The stimuli were 2 msec duration at twice the diastolic threshold value. The coupled electrical stimulation was performed with a Medtronic model 5837 stimulator. Coupling to P waves was done at decreasing intervals. When tachycardia was induced, it was stopped by an appropriately coupled stimulus.

**Terminology**

The different intervals were determined as follows. PS: beginning of spontaneous sinus depolarization to stimulus deflection (coupling interval in the figure). SP-: stimulus deflection to intrinsicoid deflection of the endocavitary atriogram: atrial response delay to stimulation. SH-: stimulus deflection to first rapid deflection of the His bundle activity: atrium to His conduction time. SR- : stimulus deflection to beginning of the delta wave: stimulus propagation to the ventricles. SJ- : stimulus deflection to end of the longest ventriculogram: total heart depolarization time. All intervals are expressed in milliseconds (msec).

**Results**

**Right atrial stimulation** Fig. 3 and 4 illustrate the effect of electrically induced atrial extrasystoles at decreasing PS. From

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1 U.S.C.I. 5651 6F.
2 Elecath 6F.
3 Elema Schönander Mingograph model EM 34.
PS of 640 msec to PS of 310 msec in Case 1 and from PS of 700 msec to PS of 420 msec in Case 2, there is a constant SR⁻ interval (105 and 140 msec, respectively). Conduction from stimulus to beginning of ventricular depolarization is therefore constant. With further shortening of the coupling interval, SR⁻ increases; this is due to an intra-atrial delay as shown by a roughly parallel increase of SP⁻. On the other hand, there seems to be a relation between the increasing delay of the SH⁻ and the increasing total depolarization time. In the actual tracing of Case 1 shown in Fig. 5, the His deflection is after the beginning of QRS. With a shorter PS, His deflection progresses in the QRS. In addition, as compared to sinus beats, the pre-excitation appearance is augmented. In Case 2, rapid right atrial stimulation (Fig. 6) induces the same effects as coupled RA stimulation: lengthening of SJ⁻ and constant SR⁻. LA stimulation is shown for comparison (see later results).

Left atrial stimulation Fig. 7 to 10 illustrate the effect of electrically induced left atrial extrasystoles. As for right atrium, the SR⁻ is constant (90 msec and 100 msec) but shorter than in right atrial stimulation and varies only when there is an intra-atrial delay (increasing SP⁻). Unlike right atrial stimulation, SJ⁻ is constant. Similarly, the His deflection, identified only in Case 1, came after the beginning of the QRS. The SH⁻ interval seems to be shorter than in right atrial stimulation: 125 msec versus 150 msec for a PS of 550 msec. In Case 2, rapid left atrial stimulation induces the same effects as coupled left atrial stimulation: constant and shorter SR⁻ and constant SJ⁻ (Fig. 6).

In Fig. 11 and 12, the constancy of SJ⁻ in left atrial stimulation is re-emphasized and contrasted to increasing SJ⁻ in right atrial stimulation. The evolution of SH⁻ is also illustrated.

Retrograde conduction Typical episodes of tachycardia in Case 1 (Fig. 13) were triggered by an atrial extrasystole at short coupling interval (PS: 200 msec for RA and 340

FIG. 6 Case 2. Rapid right atrial and left atrial stimulation. At increasing rate, consider the increasing SJ⁻ interval in right atrium as opposed to constant SJ⁻ in the left atrium.

FIG. 7 Case 1. Coupled left atrial stimulation. To read the ordinate, add 100 for SH⁻ and 200 for SJ⁻. Note the constant SR⁻ measuring 80 msec (shorter than in the right atrium 105 msec). The SJ⁻ is also constant at 210 msec. Two PH⁻ could be measured at 125 msec.
His bundle recording and electrical stimulation of atria

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The premature beat is then conducted with a longer PR and is preceded by a His potential. The pre-excitation aspect disappears. This extrasystole initiates a reciprocating tachycardia with normalized QRS and an H wave in front of each QRS. Moreover the analysis of the retrograde conduction to the atria shows that the depolarization of the high left atrial (HLA) precedes the depolarization of the low right atrial (LRA) at the tricuspid level by 30 msec. The same mechanism of tachycardia was noted in Case 2 (Fig. 14). The degree of preactivation of the left atrium is less but real as witnessed by the synchronism of the intrinsicoid deflection recorded in the low right atrium and the high left atrium.

Discussion

In the WPW syndrome, the normal His bundle activation is short-circuited, as exemplified by a His bundle deflection contemporary with or later than the delta wave. The early activation of the ventricles is, therefore, not dependent on the activation wave coming from this His depolarization. The ventricles are preexcited by an anomalous activation wave coming from a pathway that conducts faster than the normal pathway.

As we shortened the coupling intervals of the right atrial stimulation, the PR remained constant. The impulsion from the atrium to the ventricles travelled at a constant speed through the accessory fibres. However, there was an increasing delay in the normal conduction pathways as expected in normal AV nodal conduction. This produced the disappearance of the His bundle deflection into the ventriculogram. Parallel to the increasing

![Graph](https://example.com/graph.png)

**FIG. 8** Case 2. Coupled left atrial stimulation. To read the ordinate, add 200 for SJ. Again note the constancy of the intervals SR and SJ. They increase only when there is an intra-atrial delay, at PS 370 msec.

![Graph](https://example.com/graph2.png)

**FIG. 9** Case 1. Coupled left atrial stimulation. With shorter PS, SJ is constant. His potential (H) is visible only in the first two coupled beats. Compare the pre-excitation, the shorter SR, and the constant SJ to Fig. 5.
**FIG. 10** Case 2. Coupled left atrial stimulation. No H− visible in the coupled beats. Note again the short SR− and the constant SJ−.

**FIG. 11** Case 1. Comparison of SJ− and SH− intervals in the right atrium versus left stimulation. To read the ordinate, add 100 for SJ−. The increasing SJ− in parallel to SH− in right atrial stimulation is well delineated as opposed to the constancy of the same intervals in left atrial stimulation.

**FIG. 12** Case 2. Comparison of SJ− and SH− intervals in the right atrium versus left atrial stimulation. To read the ordinate, add 100 for SJ−. Increasing SJ− right atrium, constant SJ− in left atrium.
there was also an increasing SJ-. This is due to the fact that the latter part of ventricular activation is dependent on normal AV conduction. A delay in this conduction will result in a delay in ventricular activation and thus an increasing total depolarization time. The increase in the pre-excitation aspect is explained by the more and more important participation of the accessory conduction pathway to cardiac activation.

In left atrial stimulation, the pre-excitation started earlier as if the stimulation was delivered almost directly to the anomalous conduction fibres. This was responsible for the shorter SR- interval. The SJ- remained constant witnessing that all ventricular depolarization was taken in charge by the excitation coming from the anomalous pathway.

It looks to us as if, at long coupling intervals, the interatrial conduction delay represented by the difference between the SR- intervals as measured from right and left atrial excitation leaves sufficient time for normal His bundle conduction to contribute to the ventricular depolarization when the activation originates from the right atrium. This also implies that the stimulus must reach the left atrium to activate the accessory fibres. When the pre-excitation pathway is directly stimulated in the left atrium, there is no time for the impulse travelling through the His bundle to take part in ventricular depolarization. Activation is then totally accomplished by the anomalous pathway.

Another finding is a shorter SH- interval induced by left atrial stimulation in the first patient: the position of the electrodes in the atria is not the cause. A difference in refractory period in the AV node could not be eliminated. A possible explanation is represented by the existence of fibres coming from the anomalous pathway to the AV node (preferential intra-auricular conduction). In fact this pre-excitation of the normal His bundle pathway is difficult to reconcile with an exclusive activation of the ventricles by the accessory pathway. It must be assumed that the impulse coming prematurely to the common bundle is more slowly conducted and then blocked downstream by a refractory state. At shorter coupling intervals SH- lengthens due to the augmentation of the intranodal delay and H disappears in the QRS.

Regarding the tachycardia, it was our experience, in common with others (Wellens et al., 1971), that the initiation and termination of tachycardia in these two patients were easier from the left cavities particularly from the left atrium. In our two cases, the pattern was the one already described by others.
movement is initiated. The anterograde conduction from the atria to the ventricles proceeds through the normal bundle of His system as shown by the His deflection in front of the normalized QRS complexes. Retrograde conduction is probably via the anomalous bundle. In addition, retrograde conduction to the atria was faster to the left than to the right, a fact also demonstrated by Wellens et al. (1971).

The left atrial stimulation showing a shorter PR', a constant SJ', the facility of inducing tachycardia in the left cavities, the faster retrograde conduction to the left atrium are all arguments in support of the fact that in the WPW syndrome type A the anomalous pathway lies preferentially on the left side of the heart.

References


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