Role of Retrograde His Purkinje Block in the Initiation of Supraventricular Tachycardia by Ventricular Premature Stimulation in the Wolff-Parkinson-White Syndrome

MASOOD AKHTAR, MOHAMMAD SHENASA, and DONALD H. SCHMIDT,

University of Wisconsin Mount Sinai Medical Center,
Milwaukee, Wisconsin 53201

ABSTRACT The precise mechanisms for paroxysmal reentrant supraventricular tachycardia (PSVT) initiation during right ventricular premature stimulation (V2 method) were analyzed in 14 consecutive patients with Wolff-Parkinson-White Syndrome in whom the PSVT was inducible during retrograde refractory period studies. 9 patients had left-sided and the remaining 5 of 14 had right-sided ventriculo-atrial (VA) accessory pathway (AP). At the basic cycle lengths (V1V1) ranging from 550 to 900 ms (mean, 657.1±139.5), closely coupled V2 (mean V1V2, 357.3±59.2 ms, range 320–500) produced retrograde His bundle (H2) activation via the bundle branches and retrograde atrial (A2) activation via the AP. As the V1V2 were further shortened, the V2 showed a retrograde block in the His Purkinje system (HPS) and conducted to the atria via AP in 9 of 14 cases. Subsequently, the A2 impulse conducted anterograde over the atrioventricular node-HPS to initiate a PSVT or an atrial echo response in all nine cases. In none of the patients was a PSVT induced by V2 when the latter produced retrograde H2 activation via the bundle branches. In 10 of 14 cases, however, the retrograde H2 was followed by a V3, due to macroreentry in the HPS. The V3 in turn blocked retrogradely in the HPS while producing A3 via the AP to initiate a PSVT or an atrial echo response in 9 of 10 cases. Retrograde block of V2 and/or V3 in the HPS resulted in PSVT initiation in 13 of 14 cases, whereas in the remaining 1 case the exact mechanism was not clear. In none of the patients in this series was the PSVT initiated with a retrograde block of V2 in the atrioventricular node with or without concomitant retrograde A2 activation via the AP. We conclude that within the ranges of cycle lengths tested, a retrograde block of V2 and/or V3 in the HPS is the most common mechanism for initiation of PSVT during ventricular premature stimulation in patients with the Wolff-Parkinson-White Syndrome.

INTRODUCTION

Paroxysmal reentrant supraventricular tachycardia (PSVT),\(^1\) utilizing atrioventricular (AV) node and His Purkinje system (HPS) anterogradely, and an accessory pathway (AP) in the retrograde direction, is relatively common in patients with the Wolff-Parkinson-White (WPW) Syndrome (1–4). Several studies have demonstrated that PSVT in these patients can be initiated with atrial as well as ventricular premature beats. It has also been previously demonstrated that during ventricular premature stimulation, the ability or inability to induce PSVT depends upon the various patterns of ventriculo-atrial (VA) conduction along the normal and the AP (5–8). Although addressing themselves to the VA conduction along the normal pathway in patients with the WPW Syndrome, studies to date have not made a clear distinction between conduction along the HPS vs. the AV node, the two components of normal pathway. Since the HPS and AV node have different electrophysiological properties and therapeutic responses, a clear understanding of the role of retrograde conduction in

\(^1\)Abbreviations used in this paper: Ae, atrial echo; AP, accessory pathway; AV, atrioventricular; ERP, effective refractory period; HPS, His Purkinje system; PSVT, paroxysmal reentrant supraventricular tachycardia; VA, ventriculo-atrial; WPW, Wolff-Parkinson-White.

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the HPS and the AV node in the initiation of PSVT is desirable. This report demonstrates the role of retrograde block within the HPS in the initiation of PSVT during right ventricular premature stimulation in 14 patients with the WPW Syndrome.

METHODS

Right heart catheterization was performed in a post absorptive, nonseparated state, using local anesthesia. The nature of the procedure was explained to all patients and signed consents were obtained. Quadripolar electrode catheters were percutaneously inserted into peripheral veins and positioned in the region of the high right atrium, AV junction, coronary sinus, and right ventricle. The catheters were used for local bipolar electromgram recordings and/or electrical stimulation using techniques previously described (9). Surface electrocardiogram leads I, II, and VI, intracardiac electrograms, and time lines were simultaneously displayed on a multichannel oscilloscope and recorded on a frequency-modulated tape. The recordings were subsequently retrieved for analysis at a paper speed of 100–150 mm/s. Intracardiac stimulation was performed using digital stimulator capable of delivering rectangular impulses with adjustable voltage and duration. During the conduct of these studies, patient isolation existed and all equipment was carefully grounded.

Retrograde refractory period studies were performed using ventricular extrastimulus method. Premature stimuli were introduced after eight ventricular paced beats of predetermined constant cycle length. The coupling intervals were gradually reduced by 10–20 ms until ventricular muscle retractoriness was encountered.

Definition of terms. A complete set of definitions for anterograde and retrograde conduction and refractory period studies has been previously published (9). Pertinent definitions repeatedly used in this text will be outlined here.

The A-H interval was measured from the onset of low atrial electrogram to the beginning of His bundle potential. The H-V and RB-V intervals were measured from the onset of His and right bundle branch deflections to the earliest detectable ventricular activity, either on the surface electrocardiogram or the local intracardiac tracings.

Atrial echo (AE) response represents spontaneous retrograde excitation of the atrium exclusively via the accessory pathway occurring during nonpaced beats.

For an easier comprehension some of the terms used and electrophysiologic parameters measured during retrograde refractory period studies are graphically depicted in Fig. 1. The S, V, and H represent stimulus artifact, ventricular, and retrograde atrial deflections, respectively, during the basic drive, whereas S, V, H, and A represent stimulus artifact, ventricular, His bundle, and atrial deflections following the premature beat.

The retrograde activation in response to V2 of the H2 via the bundle branches and the A2 via AP is shown in panel A. Exclusive VA conduction via the AP during both V1 and V2 is suggested by the sequence of atrial activation and its lack of change following V3 despite significant delay along the normal pathway (V1H2 interval).

The effective refractory period (ERP) of AP (panel B) is the longest V1V2 where V2 blocks in the AP.

The ERP-AV node (panel C) is the longest V1H2 where H2 is not followed by A2. V1H2 is taken in lieu of retrograde H,H2 since H1 is generally not visible during the basic drive, and it is assumed that V1H2 interval is constant (9). It should be noted that ERP-AV node can only be determined after retrograde ERP of AP pathway is encountered.

ERP-HPS (panel D) is longest V1V2 where V2 blocks below the His bundle recording site (i.e., HPS).

RESULTS

Anterograde and retrograde conduction and refractory period studies were performed in 22 consecutive patients with WPW Syndrome who had symptomatic PSVT. In only 14 of 22 patients was the PSVT inducible during right ventricular premature stimulation, and findings in these 14 cases form the basis of this report. In the remaining 8 of 22 cases, PSVT could not be elicited during ventricular premature stimulation for the reasons previously published (7). 12 of the 14 had ventricular preexcitation during sinus rhythm, whereas the remaining two (patients 5 and 8) had electrocardiographically concealed left sided AP. The pertinent clinical and electrophysiologic data are summarized in Table I. At the time of study all were in sinus rhythm and were not taking any cardioactive medications. Two patients had associated arteriosclerotic heart disease, and another two had mitral valve prolapse, whereas the remaining cases did not have clinically detectable structural heart disease. Although two or more basic ventricular cycle lengths were scanned in 10 of 14 patients, to simplify presentation of the results only the data from the shortest available cycle lengths in each case will be presented in detail.

Retrograde refractory period studies. Ventricular premature stimulation was initiated at V1V2 intervals which measured 50–100 ms less than the basic drive, and the coupling intervals were progressively shortened. During the basic drive (V1) six patients (patients 2, 3, 5, 7, 10, and 13, and Table I) demonstrated retrograde atrial depolarization exclusively via the AP, whereas fusion retrograde atrial activation was observed in the others (10). In the latter cases a change from fusion atrial activation to exclusive activation via the AP was noted as the V1V2 were further shortened. This change in retrograde atrial activation was noted at V1V2 intervals which had not yet produced emergence of a His deflection from the V2 electrogram. At the basic cycle length range of 550–900 ms (Table I), the His bundle deflection first emerged from V2 at V1V2 intervals ranging from 320–500 ms. Since in patients with retrogradely functioning AP the His bundle deflection following V2 could result from either retrograde conduction via the bundle branches or antegrade depolarization via the AV node following A2 activation over the AP, the behavior of His bundle activation was closely analyzed (11). In all nine patients with left-sided (cases 1–9, Table I) and 2 of 5 cases (patients 11 and 12) with right-sided VA AP, the separation of retrograde A2 from V2 deflection (on the His bundle electrogram tracing) was sufficient to provide a clear segment where the presence of a His deflection could be identified. In all
of these 11 patients the His bundle deflection preceded the retrograde A₂ deflection (Figs. 2 and 3) during its initial appearance, indicating retrograde His bundle activation via the bundle branches (H₂); the designation of H₂ henceforth will imply such an occurrence. With further shortening of the V₁V₂ intervals, the V₂H₂ delays showed almost linear increases as long as the retrograde H₂ deflection was still identifiable and preceded or coincided with the retrograde A₂ electrogram (Figs. 2 and 3). This relationship between V₁V₂ and V₂H₂ intervals is typical of retrograde H₂ activation via the bundle branches and has been previously demonstrated in patients without the WPW Syndrome (9, 12–15).

At the V₁V₂ intervals that produced emergence of retrograde H₂ deflection, the V₂A₂ intervals via the AP showed no further prolongation once exclusive V₂A₂ conduction via the AP was previously established (Figs. 2–4) at longer coupling intervals. Upon further shortening of the ventricular coupling intervals in 8 of

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### Table I

**Clinical and Electrophysiological Data**

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Abbreviations used in this table: ASHD, arteriosclerotic heart disease; AVN, AV node; B, block; MVP, mitral valve prolapse; NP, normal pathway; Ret, retrograde; VCL, ventricular cycle length; Vent, ventricle. L and R indicate left or right VA AP. + indicates the presence or occurrence of an event.

* ΔVH represents a change of H activation from retrograde via bundle branches to anterograde via the AV node.

† Electrophysiologic events that coincided with initiation of PSVT.

‡ Events coinciding with occurrence of a single echo beat.
these 11 cases (cases 1–7 and 11, Table I), the His deflection suddenly shifted from a location preceding or coinciding with the A₂ (on HB electrocardiogram) to a position following the A₂ (Figs. 2–4). This change in location of His bundle deflection was accompanied by an increase in V-H interval ranging from 40 to 155 msec (mean, 121.5±29.1) for a 10–30 ms (mean, 16.2±7.4) decrease in the V₁V₂ interval (Table I). The succeeding A₂H intervals in most cases measured equal to or were slightly longer than (i.e., 10–20 ms) the A-H intervals of sinus beats (except in patient 11) but significantly shorter as compared with the A-H intervals during PSVT. In all of these eight cases ventricular activation followed the His bundle activation, which in turn was followed by retrograde atrial activation via the AP (atrial echo beat or Ae), and PSVT was initiated (Figs. 2–4) in all except one case (case 11). In case 11 the Ae beat anterogradely blocked in the AV node and no sustained PSVT occurred. The sudden increase in the V-H interval was not due to retrograde block in the right bundle branch and subsequent retrograde His activation via the left bundle branch. The events can be best

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Figure 2. (Patient 3) Basic ventricular cycle length is constant at 600 ms (panels A–D). The V₁A₁ and V₁A₂ intervals measure the same in all panels despite progressive shortening in the V₁V₂ intervals. In addition to the two HB tracings, a recording from the RB was also obtained with the H-RB interval measuring a 30 ms during sinus rhythm. The V₂ in panel A depolarizes the H₂ via the left bundle branch (LBB) as suggested by H₂-RB₂ interval of 10 ms. A 10-ms shortening in V₁V₂ interval (panel B) results in an increase in the V-H from 170 ms in panel A to 225 ms in panel B, and PSVT is initiated. Note that the His deflection in panel B follows A₂, and H-RB interval equals that of sinus beats (30 ms) indicating His bundle activation via the AV node. The A₁H interval equals sinus AH (not labeled) and is significantly shorter compared with Ae-H of PSVT, suggesting lack of retrograde penetration of the AV node by V₂ impulse. The QRS following His deflection shows an LBB block (B) pattern, indicating incomplete recovery of LBB following V₂ despite retrograde bilateral block HPS. At shorter V₁V₂ of 300 ms (panel C) V₁H₂ conduction via the LBB resumes (retrograde gap HPS) and PSVT is not induced. Retrograde H₃ activation via the LBB continues in panel D at a shorter coupling interval, and a V₃ due to macroreentry HPS is noted. The V₃ produces A₃ response via the AP blocks in the HPS, and a PSVT is started. 1, 2, and V₁ surface ECG leads; RA, right atrial electrogram; RB, right bundle electrogram tracing; T, time lines.
explained as follows: The V₂ retrogradely blocked bilaterally in the HPS (i.e., no H₃), yet reached the atrium via the AP. Since the AV node was not penetrated by the V₂ impulse, the A₂ response was followed by relatively fast anterograde AV nodal conduction, i.e., A₃H interval. The longer H-V intervals and the aberrant conduction of the ensuing QRS complex noted in some cases was due to retrograde concealed conduction of V₂ in the HPS. The reasons for the above assumption, which was directly documented in two cases by simultaneous His and right bundle (RB) recordings (Fig. 2), will be discussed later. To simplify presentation of the results, the designation V₂H will imply anterograde depolarization of the His potential via the AV node following retrograde A₂ activation over the AP. The occurrence of these electrophysiologic events in individual cases, i.e., retrograde block in the HPS and a shift in His activation from retrograde via the bundle branches to anterograde via the AV node and initiation of PSVT, is depicted in the table. In the remaining 3 of 11 cases (8, 9, and 12), a retrograde block in the HPS was not noted and V₂ continued to activate H₂ via the bundle branches, as indicated by the following: (a) persistent linear relationship between V₂H and V₁V₂, (b) the His deflection never showed a shift to a position succeeding the A₂ electrogram. None of these three cases demon-
ever, a sudden increase in the V-H of 75 ms with a 20 ms decrease in $V_1V_2$ coincided with initiation of PSVT, suggesting that the initial His activation was retrograde, whereas the onset of PSVT heralded a retrograde block in the HPS, findings similar to all other cases where $V_2$ initiated PSVT.

In the one remaining patient (13), at no time did the His deflection precede $A_2$, and a change from retrograde to anterograde depolarization of the His bundle could not be documented. Although a PSVT was initiated in this case by $V_2$, discernment of the exact mechanism was not certain.

Retrograde gap phenomenon HPS (16). Following the initial retrograde block of $V_2$ in the HPS, further shortening in $V_1V_2$ intervals resulted in resumption of retrograde conduction to the His bundle (i.e., $V_2H_2$) in seven of nine cases (1, 3–7, and 11, Table I retrograde gap phenomenon HPS), as was suggested by return of $H_2$ deflection preceding or coinciding with the $A_2$ electrogram (Figs. 3 and 4). The $V_2H_2$ values following resumption of conduction were in line with those expected from a reciprocal relationship between $V_2V_3$ and $V_2H_3$ intervals and significantly shorter than the $V_2H$ values noted at longer coupling intervals (Fig. 4). Return of $V_2H_2$ conduction occurred while retrograde $A_3$ activation persisted via the AP in all cases except one (point No. 5, Fig. 3), where a concomitant retrograde block was noted along the AP. Whenever retrograde $H_2$ followed $V_2$ with or without associated $A_2$ activation via the AP, the $Ae$ or PSVT was not initiated by $V_2$ either before or after the gap zone in any of the cases in this series.

Macrobeentry HPS (12, 13). 10 of the 14 cases (3–9, 11, 13, and 14, Table I) demonstrated occurrence of macroreentry HPS in the form of $V_3$ following retrograde $H_2$ activation. The $V_3$ in turn showed a retrograde block in the HPS along the normal pathway (i.e., $H_3$) in all patients (Fig. 3). The $V_3$, however, conducted to the atrium ($A_3$) via the AP in all cases and an Ae or PSVT was started in 9 of 10 (Table I, Fig. 3). In the remaining one case the $A_3$ blocked in the AV node, and no Ae or PSVT was noted.

Findings at other basic cycle lengths. In 10 of 14 cases, two or more basic cycle lengths were tested, and the results were similar to those presented above, i.e., the PSVT was only initiated when $V_2$ and/or $V_3$ retrogradely blocked in the HPS while still activating atrial via the AP.

**DISCUSSION**

Results of this study indicate that retrograde HPS block of $V_2$ and/or $V_3$ in the HPS is a relatively common mechanism by which PSVT is initiated during right ventricular premature stimulation in patients with the WPW Syndrome (13 of 14 cases in this series). Although...
the induced V₂ may initiate PSVT by other mechanisms, present data suggest that other mechanisms are less commonly responsible for PSVT initiation during ventricular premature stimulation. In a series with 139 cases of WPW Syndrome reported by Wellens (6) it was noted that, during ventricular premature stimulation, a PSVT was most commonly initiated when V₂ retrogradely blocked along the normal pathway (30 of 139 cases) or by V₃ (9 of 139 cases) when the latter was inducible, and by other mechanisms in only 11 of 139 cases. The exact site of retrograde block (HPS vs. AV node) of V₂ or V₃ was not demonstrated, and it cannot be stated with certainty whether the findings in the report by Wellens and the present study are comparable. A recent study showed that in patients with ventricular preexcitation, programmed ventricular extrastimuli did not initiate PSVT if there was no VA conduction and there was persistent retrograde atrial fusion activation via both the AP and normal pathway (7). A narrow QRS PSVT could be elicited with V₂ when VA conduction was intact, V₂A₂ conduction occurred exclusively via the AP, and normal pathway was less deeply penetrated (7). In that report it was indicated, however, that the level or site of retrograde block along the normal pathway could not be determined.

Regular narrow QRS complex PSVT is the most common arrhythmia elicited by programmed atrial as well as ventricular premature stimulation in patients with the WPW Syndrome (3, 4). During premature atrial stimulation (A₂), the premature impulse blocks in the AP, activates the ventricles over the AV node-HPS, and returns to the atria via the AP to start the PSVT. Induction of the same PSVT with V₂, on the other hand, will be more likely if V₂A₂ conduction occurred via the AP and subsequent anterograde propagation took place along the AV node-HPS. The latter occurrence will obviously be facilitated by a retrograde block of V₂ in the normal pathway. A retrograde block of V₂ above the His bundle (AV node), however, may not permit initiation of PSVT even if accompanied by concomitant retrograde A₂ activation via the AP, because of effect of retrograde concealed conduction in the AV node. Under the following circumstances, however, V₂ may start PSVT despite AV nodal penetration (a) when V₂ produces only a partial penetration of the AV node with resultant rapid recovery of excitability, (b) initiates intranodal reciprocation, (c) conducts with long V₂A₂ via AP sufficient to allow recovery of AV nodal excitability. A retrograde block of V₂ in the HPS, on the other hand, would permit unhindered progress of A₂ impulse through the AV node, since the prior AV nodal depolarization is during the last V₁.

A support for the preceding hypothesis is provided in the present study by demonstration that the PSVT initiation did coincide with the occurrence of retrograde block of V₂ in the HPS. That the site of retrograde block of V₂ was indeed in the HPS rather than the AV node is suggested by the following observations: (A) A sudden increase in the V-H values (a mean increase of 121.8 ms from V₂H₁ to V₂H₂ for a 16.2 ms mean decrease) in V₁V₂ intervals, a behavior unlike that of retrograde H₂ activation via the bundle branches and therefore suggesting a shift from retrograde to anterograde activation of the His bundle (Figs. 2–4). (B) A return of H₂ deflection at shorter V₁V₂, producing V₂H₂ values less than V₂H₁ values (retrograde gap phenomenon HPS) noted at longer V₁V₂, and concomitant abolition of PSVT (Figs. 3 and 4).

(C) Direct demonstration of the shift from a retrograde to anterograde activation of the His bundle when both His and RB recordings were available (two cases, Fig. 2). (D) The occurrence of macroreentry HPS after retrograde H₂ activation and its disappearance despite longer V-H intervals, coinciding with the shift of His deflection to a location succeeding the A₂ (Fig. 3). (E) A consistent relationship between the A₂ and the succeeding His deflection with values approximating the sinus A-H values. This suggests that the occurrence of His deflection after A₂ was not an independent phenomenon (retrograde via bundle branches) but its occurrence was directly related to and in fact dependent upon the preceding A₂ activation (Figs. 2 and 3).

It seems reasonable to conclude, therefore, that the His deflection following the A₂ was due to anterograde depolarization via the AV node. Despite the above reasoning, questions can be raised as to whether the V₂ might still have blocked in the AV node rather than HPS, and whether the H₂ deflection was either not identifiable for some reason or was obscured by the local V₂ electrogram. These questions can be dismissed for the following reasons: (a) Once the retrograde H₂ deflection emerges, the V₂H₂ intervals generally lengthen (rather than shorten) at shorter V₁V₂ (9, 12, 14, 15). (b) The occurrence of gap phenomenon in most cases suggests retrograde H₂ was recordable whenever the His bundle was depolarized. (c) At the V₂H interval ranges noted in these cases, the His bundle activation will have to occur twice (first retrogradely and then anterogradely) within a period of 300 ms, an unlikely occurrence. In the event the V₂ retrogradely penetrates the AV node, the subsequent A₂H intervals will be expected to be much longer than the sinus beats (retrograde concealed conduction in the AV node). Such a mechanism of PSVT initiation was not noted in the present series but has been previously described, and may operate more frequently at shorter cycle lengths due to abbreviation in the refractoriness of the His. For similar reasons a retrograde AV nodal block of ventricular impulse may also
be a more common mechanism of PSVT initiation during rapid incremental ventricular pacing rather than premature stimulation (8).

It was repeatedly observed during this study that whenever V₂ produced a retrograde H₂ response, the PSVT could not be initiated. Since concomitant retrograde A₂ activation in most instances took place exclusively via the AP, the collision of the two impulses via the normal pathway and AP must have occurred in the AV node rather than the atrium (7, 17). Retrograde penetration and concealed conduction in the AV node was most probably responsible for preventing the forward progress of the A₂ impulse. It is easily understandable, therefore, why the onset of retrograde block in the HPS coincided with initiation of a PSVT or an Ae response. It may also be pointed out that the response of HPS and AP to V₂ was not significantly different between right- and left-sided AP, aside from the fact that the location of A₂ on the two electrograms in patients with right VA AP made analysis of events more difficult.

When an H₂ response followed V₂, the PSVT only occurred if V₂ followed. Although in this group of patients the V₂ phenomenon occurred due to macroentry HPS, a V₂ from other mechanisms—local reentry, etc.—will be expected to produce the same results (13). The initiation of an Ae response or PSVT following V₂ was also facilitated by a retrograde block of V₂ in the HPS and concomitant retrograde A₂ activation via the AP, as was noted in 9 of 14 cases in this series.

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REFERENCES


