Limitation of tachycardia zone resulting from longitudinal dissociation of the atrioventricular node in concealed pre-excitation

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SUMMARY Two cases with a concealed left-sided accessory atrioventricular bypass tract are described. In both, functional longitudinal dissociation of the atrioventricular node narrowed the range of atrial premature beat coupling intervals which could initiate re-entry using the accessory pathway. In case 1 early premature atrial beats were followed by an atrioventricular nodal re-entrant echo. The atrial echo pre-empted retrograde conduction over the Kent bundle and thus limited the development of paroxysmal supraventricular tachycardia. In case 2 atrioventricular nodal conduction showed typical features ascribed to dual atrioventricular nodal pathways. In addition there was a bradycardia-related retrograde block in the concealed accessory pathway. Early premature atrial beats, because of exclusive “slow pathway” anterograde conduction, arrived at the ventricles during the period of bradycardia-dependent retrograde block and failed to initiate a macro re-entrant tachycardia. This study shows that (1) longitudinal dissociation within the atrioventricular node may limit the ability to initiate tachycardia in patients with concealed pre-excitation; and (2) discontinuous atrioventricular nodal conduction curves occasionally help to reveal bradycardia-related retrograde block in a concealed accessory pathway.

Discontinuous atrioventricular nodal conduction curves and atrioventricular nodal re-entry are common electrophysiological responses to premature atrial stimulation in man.1 The former are believed to reflect dual atrioventricular nodal pathways and the latter dual pathways, functional longitudinal dissociation, or reflection within the atrioventricular node.1 2 These phenomena have recently been described in a few patients who also had manifest3-5 or concealed pre-excitation.6-9 In most of these reported cases the atrioventricular nodal conduction characteristics were unrelated to the initiation or maintenance of the paroxysmal supraventricular tachycardia.6-8 In some, the critical atrioventricular conduction delay required for macro re-entry was achieved by block in the fast atrioventricular nodal pathway, and thus initiation of the tachycardia was enhanced by “duality” of atrioventricular conduction.5 9 In a few patients with an accessory pathway, atrioventricular nodal re-entry itself proved to be the mechanism of the tachycardia.5 8 10

The purpose of the following report is (1) to present two unusual cases of concealed pre-excitation where atrioventricular nodal re-entry and dual atrioventricular nodal conduction limited rather than enhanced initiation of paroxysmal supraventricular tachycardia with early atrial extrasystoles, and (2) to show for the first time, a bradycardia-related retrograde block in a concealed accessory pathway.

Case reports

Case 1, a 57-year-old man, and case 2, a 54-year-old woman, were studied because of long histories of paroxysmal supraventricular tachycardia. Electrophysiologival evaluation of the patients included intracardiac recordings from several sites and incremental pacing and extrastimulation.6 9 11 The two cases had the following features in common. (1) Routine electrocardiograms were normal. (2) Ventricular pre-excitation was not produced by right or left atrial pacing and extrastimuli. (3) The presence of a left-sided accessory atrioventricular pathway capable of retrograde conduction only could be proved by ventriculoatrial conduction studies and endocardial atrial mapping.6 11 (4) The concealed accessory atrioventricular pathway was responsible for the initiation of atrial fibrillation.

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Tachycardia zone in concealed pre-excitation

Fig. 1 (case 1)  Limitation of the tachycardia zone by atrioventricular nodal re-entry. Left atrial pacing from the coronary sinus is performed at a basic cycle length of 600 ms. In A, a non-sustained supraventricular tachycardia resulting from macro re-entry over a left-sided atrioventricular bypass tract is initiated by a premature atrial impulse with coupling time of 320 ms. Note the left-to-right activation sequence of the atrial echo beats (Ae). In B, the coupling time is 310 ms. After prolongation of the A2-H2 interval to 160 ms, there is a single atrioventricular nodal re-entrant complex, and macro re-entry does not occur. Note the simultaneous atrial and ventricular activation and the low-to-high right atrial activation sequence of the echo in B. Panel C shows that blocked premature stimuli are not followed by atrial echoes.

Each panel is arranged from top to bottom as follows: electrocardiographic lead III, chest lead I (V1), high right atrial lead (HRA), His bundle lead (HBE), and coronary sinus lead (CS). S1, A1, and H1 are, respectively, the stimulus artefact, low right atrial deflection, and His deflection of the last basic drive beat; S2, A2, and H2 are the respective deflections of the atrial extra stimulus.

Paper speed in each illustration is 100 mm/s. All numbers in this and subsequent figures are in ms.
for the paroxysmal supraventricular tachycardia in both patients. 

Functional longitudinal dissociation of the atrioventricular node with early atrial extrasystoles limited the potential tachycardia zone with regard to initiation of atrioventricular macro re-entry.

In case 1, evidence for participation in the paroxysmal supraventricular tachycardia of a left-sided accessory pathway included the following. 

(1) The earliest atrial activation during the tachycardia was recorded at the mid-coronary sinus, and this preceded the low right atrial electrogram in the His bundle lead by 40 ms (Fig. 1A and 2A). 

(2) Ventriculoatrial conduction times during tachycardia were 50 ms longer on beats with left bundle-branch block than on beats with normal intraventricular conduction (Fig. 2). 

In case 2, the following data supported macro re-entry over a concealed left-sided accessory tract.

(1) The left atrial electrogram recorded at the distal coronary sinus during the tachycardia preceded the low right atrial electrogram by 50 ms (Fig. 3B and 4). 

(2) Single extrasystoles delivered at the right ventricular apex during tachycardia pre-excited the left atrium by up to 60 ms. This occurred at a time when the ventricular extrasystole must have found the normal atioventricular node—His bundle pathway refractory from anterograde conduction of the previous beat (Fig. 4).

Analysis of the range of A1–A2 intervals producing tachycardia in case 1 disclosed the following (Fig. 1).
Fig. 3 (case 2)  Limitation of the tachycardia zone because of a sudden “jump” in the A-H interval of an early premature beat. Right atrial pacing is performed at a basic cycle length of 500 ms. In B, supraventricular tachycardia using a concealed left-sided accessory pathway is initiated. With a coupling time 10 ms longer (A), the critical atrioventricular delay for re-entry is not yet achieved. With a coupling time 10 ms shorter (C), there is a sudden jump in the A1-H1 interval and again there is no tachycardia. Abbreviations are the same as in Fig. 1. CS is a distal coronary sinus lead. Numbers above V1 leads are V1–V2 intervals.
When left atrial pacing was performed at a cycle length of 600 ms, the paroxysmal supraventricular tachycardia could be induced by atrial extrasystoles over a narrow range of A1–A2 intervals (330–315 ms). The atrioventricular nodal conduction times on these premature beats measured 115 to 130 ms. When A1–A2 was decreased to 310–300 ms, the A2–H2 interval further increased (Fig. 1B). Nevertheless, these early extrastimuli failed to initiate the tachycardia. The inability of these extrasystoles to induce macro re-entry was secondary to appearance of an atrial echo which pre-empted retrograde conduction over the accessory pathway. The mechanism of the atrial echo was atrioventricular nodal re-entry rather than intra-atrial re-entry since (a) the atrial echo beat had a low-to-high right atrial activation sequence (Fig. 1B); (b) a critical AH interval of 160 ms was required for the echo to appear (Fig. 1B); and (c) the atrial echo never occurred when A2 blocked in the atrioventricular node (Fig. 1C). Significant intra-atrial delay of the premature impulses could not be held responsible for retrograde block in the accessory pathway. Therefore, it is reasonable to assume that had functional atrioventricular nodal dissociation and atrioventricular nodal re-entry not occurred, the tachycardia zone, at least at the cycle length tested, would have ranged from 330–300 ms. Atrioventricular nodal re-entry was responsible for narrowing of the tachycardia zone by 15 ms.

In case 2, a paroxysmal supraventricular tachycardia incorporating the concealed accessory pathway was initiated over a narrow range of A1–A2 intervals. With right atrial pacing at a cycle length of 500 ms, the tachycardia zone ranged from 275–260 ms (Fig. 3 and 5). By decreasing A1–A2 to 260–230 ms, there was a sudden increase in the A2–H2 intervals (Fig. 3 and 5). This was probably the result of block in a “fast” and exclusive conduction through a “slow” atrioventricular nodal pathway.1 Whenever there was very slow conduction in the atrioventricular node, the...
extrastimulus failed to initiate the tachycardia (Fig. 3C). Atrioventricular nodal re-entrant complexes were also not observed.

The only mechanism that could reasonably explain the above findings is to postulate a bradycardia-dependent retrograde block in the accessory pathway. If this were the case, early premature beats, by conducting exclusively through a slow atrioventricular nodal pathway, would reach the ventricles at a period when the concealed accessory atrioventricular pathway was unresponsive because of the bradycardia-related block. This hypothesis was tested by the ventricular extrastimulus technique. The pertinent findings are illustrated in Fig. 6. Left atrial pacing was performed at the same cycle length as during the anterograde studies. After eight basic drive beats a right ventricular extrastimole was delivered. This methodology was employed to allow the atria to recover fully after the last basic drive stimulus. Thus, retrograde conduction of the ventricular extrasystole was solely dependent on the ability of the normal and anomalous pathways to deliver the impulses to the atria. Ventricular extrasystoles late in diastole all conducted to the atria. The $V_2-A_2$ intervals progressively increased with increasing prematurity. The low right atrial electrogram ($A_2$) preceded the left atrial electrogram ($A_3$) by a constant interval, 40 ms. This suggested that retrograde conduction of late ventricular premature systoles occurred through the atrioventricular node. The shortest coupling time of $V_2$ showing this retrograde activation sequence was 430 ms (Fig. 6A). When $V_1-V_2$ was decreased to 420 ms, retrograde conduction failed to occur (Fig. 6B). The first supraventricular complex following $V_2$ in this panel is a sinus node escape. There was complete retrograde block between coupling times of 420–350 ms (Fig. 6B and 6C). With a $V_1-V_2$ of 340 ms retrograde conduction reappeared, but this time the atrial activation sequence was left to right (Fig. 6D). This suggested that retrograde conduction occurred over the accessory pathway. This type of conduction was manifest down to a $V_1-V_2$ interval of 310 ms (Fig. 6E). With a $V_1-V_2$ of 300 ms or less, there was again complete retrograde block (Fig. 6F).

The above findings can be interpreted as evidence for bradycardia-dependent retrograde block in the concealed accessory pathway. With $V_1-V_2$ intervals of 350 ms or longer, probably all premature ventricular beats blocked in the accessory pathway. The constant $A_2-LA_3$ intervals down to a $V_1-V_2$ of 430 ms and complete ventriculoatrial block between $V_1-V_2$'s of 420–350 ms favour this assumption.

At coupling
intervals shorter than 350 ms, retrograde conduction occurred exclusively over the accessory pathway until its refractory period of 300 ms was reached. These findings show a remarkable correlation with the anterograde studies. Whenever a paroxysmal supraventricular tachycardia was initiated by a premature atrial impulse, the $V_1$-$V_2$ interval was in the range where retrograde conduction could occur through the bypass tract (Fig. 3 and 5). Whenever there was exclusive "slow pathway" anterograde conduction, the $V_1$-$V_2$ intervals fell in the range of brady-cardia-dependent retrograde block. Atrioventricular re-entry was not initiated (Fig. 3C and 5). Fig. 6G illustrates that bradycardia-dependent retrograde block was a function of the basic cycle length. During tachycardia an atrial extrastimulus which caused a subsequent V-V interval of 590 ms did not result in retrograde block in contrast to long V-V intervals during regular pacing at a longer cycle length than the tachycardia. The exact ranges of retrograde accessory pathway conduction at the rate of the tachycardia were not tested by the ventricular extastimulus method.

Discussion

In the presence of a concealed anomalous bypass tract macro re-entry can be induced by premature atrial...

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Fig. 6 (case 2)  Demonstration of bradycardia-dependent retrograde block in a concealed left-sided atrioventricular accessory pathway (panels A to F). $S_1$ represents the last two stimulus artefacts of eight basic drive stimuli delivered to the coronary sinus at a cycle length of 500 ms. $S_2$ is delivered to the right ventricular apex. Numbers in the $V_1$ leads are coupling times ($V_1$-$V_2$ intervals) of the ventricular extrasystoles. $A_2$ and $LA_2$ are the low right atrial and left atrial deflections conducted from $V_2$. The ventricular extrasystoles in each panel are followed by a sinus escape except for panel E where there is a ventricular echo. The range of coupling times with retrograde conduction over the accessory pathway is from 340 to 310 ms. Panel G shows that (a) exclusive slow pathway anterograde conduction per se is not responsible for retrograde block in the accessory pathway, and (b) the bradycardia-dependent block is rate-related. See detailed explanation in the text.
Stimuli that have atrioventricular conduction times sufficient to overcome refractoriness in the accessory pathway and in the atrium. The range of coupling times of premature atrial impulses capable of initiating the tachycardia (that is the tachycardia zone) is dependent on several factors including the basic drive rate and the site of stimulation in regard to the site of the accessory pathway. Once a critical coupling time is reached, however, where the premature atrial impulse initiated paroxysmal supraventricular tachycardia, premature stimuli with shorter coupling times usually also induce the tachycardia up to a point where A2 fails to conduct to the ventricles. The only exception is the rare case when very early extra impulses result in pronounced intra-atrial delay or in a repetitive atrial response which pre-empts conduction over the concealed accessory pathway.

In the two cases described in this paper, the effects of longitudinal dissociation in the atrioventricular node limited the ability to induce the tachycardia with early premature atrial stimuli. In case 1, a non-sustained supraventricular tachycardia using an accessory pathway was initiated by premature atrial impulses which developed a critical A2-H2 interval of 115 ms or longer (Fig. 1). As the premature atrial coupling intervals shortened, atrioventricular nodal conduction lengthened. Eventually atrioventricular nodal echo occurred. This atrioventricular nodal echo prevented macro re-entry over the accessory pathway by creating refractoriness in the atrium (Fig. 1B). Since the atrial echo caused by atrioventricular nodal re-entry was too early to be conducted to the ventricles, the re-entry process was limited to one single echo and a tachycardia did not develop (Fig. 1B). Significant intra-atrial delay or repetitive atrial responses were not present with early atrial extra impulses. Therefore, it is reasonable to assume that atrioventricular nodal re-entry alone was responsible for limiting the potential tachycardia zone.

In case 2, a critical A-H interval of 140 ms was required for the atrial premature stimuli to overcome refractoriness in the retrograde pathway. At atrial coupling intervals of 260 ms or shorter, however, there was a sudden increment in the atrioventricular nodal conduction times (Fig. 5). At these shorter coupling times with long A-H intervals neither atrioventricular nodal re-entrant echoes nor re-entry over the accessory pathway were observed (Fig. 3C). Inspection of the atrioventricular nodal conduction curves of this patient (Fig. 5) disclosed the features ascribed to dual atrioventricular nodal pathways. Failure of re-entry over the accessory pathway at these long A2-H2 intervals could not be reasonably related to concealed anterograde penetration of A2 into the accessory pathway or to atrial refractoriness from the premature atrial stimulus. Inability of the early premature stimuli to initiate paroxysmal supraventricular tachycardia proved to result from a bradycardia-dependent retrograde block in the concealed accessory pathway. The "role" of dual atrioventricular conduction in this case was to delay arrival of impulses from early atrial extrasystoles at the ventricular insertion of the bypass tract and thus prevent them from returning to the atria. The inner limit of the bradycardia-dependent block in the accessory pathway proved to be 350 ms (Fig. 6C). It is reasonable to suspect that in the absence of a sudden jump in the A2-H2 intervals with short coupling times of premature stimuli the V1-V2 intervals of at least some early atrial stimuli would have remained in the responsive zone of the accessory pathway and thus would have initiated the tachycardia. The sudden jump in the A2-H2 intervals therefore may have limited the potential tachycardia zone with regard to atrioventricular re-entry.

Bradycardia-dependent accessory pathway block had been previously shown to occur in patients with overt pre-excitation. To our knowledge our case 2 is the first patient reported where a bradycardia-related retrograde block was proved in a concealed atrioventricular bypass tract. If the range of bradycardia-dependent retrograde block is "far to the right", that is retrograde block in the bypass tract occurs only late in diastole, right ventricular stimulation studies may not be very helpful in disclosing this peculiar disturbance. Because of the proximity of the normal atrioventricular pathway to the right ventricular pacing site, late coupled ventricular extrasystoles may result in atrial activation over the normal pathway even in the presence of a remote left-sided concealed pathway. Therefore, detection of a bradycardia-related block in the accessory pathway with late coupled V2's may be difficult. In this case, discontinuous atrioventricular nodal conduction curves, as in our patient, may help to show a bradycardia-related block in a concealed accessory pathway.

Previous studies have shown that in the presence of a concealed atrioventricular bypass tract a sudden increase in the atrioventricular nodal conduction time of early premature atrial stimuli can help to initiate a supraventricular tachycardia. Moreover, atrioventricular nodal re-entry may be the mechanism of the paroxysmal supraventricular tachycardia in some of these patients. Our study shows that atrioventricular nodal conduction over a slow pathway and atrioventricular nodal re-entry can also limit the potential tachycardia zone in patients with concealed accessory bypasses. Pharmacological interventions eliminating the above phenomena could paradoxically
enhance the occurrence of paroxysmal supraventricular tachycardia in these patients.

References


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Limitation of tachycardia zone resulting from longitudinal dissociation of the atrioventricular node in concealed pre-excitation.

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*Br Heart J* 1981 46: 302-310
doi: 10.1136/hrt.46.3.302

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