Mechanisms of junctional tachycardia showing ventricular pre-excitation

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SUMMARY Over a period of five years 12 patients underwent electrophysiological studies for the investigation of recurrent tachycardias which showed ventricular pre-excitation. Nine patients had a type B pattern and two a type A. One patient had episodes of both types. Dual atrioventricular nodal pathways were found in six of seven patients with atrioventricular nodal re-entrant tachycardia mechanisms. Single direct atrioventricular accessory pathways were present in four patients, single nodoventricular pathways in five, and multiple pathways in three. Twenty one tachycardias were induced, of which 13 showed ventricular pre-excitation. Five patients had nodoventricular pathway conduction during atrioventricular nodal tachycardia and one during atrioventricular re-entrant tachycardia. Only three patients had simple antidromic tachycardia and one additional atrioventricular nodal tachycardia with bystander atrioventricular accessory conduction. Three patients had three different tachycardias, three had two types, and six had one type.

Thus junctional tachycardias showing ventricular pre-excitation are often associated with multiple mechanisms and complex anatomical and functional substrates. An accessory pathway was an essential component in only six of 13 tachycardias showing ventricular pre-excitation. Determination of the tachycardia mechanism requires detailed study and analysis.

Several mechanisms of tachycardia have been demonstrated in patients with the Wolff-Parkinson-White syndrome.1 By far the commonest mechanism is atrioventricular re-entrant tachycardia with retrograde conduction over the accessory atrioventricular connexon and anterograde conduction over the atrioventricular nodal-His bundle pathway. It is well known, however, that tachycardia may also be caused by re-entrant activity in the reverse direction producing a tachycardia with broad "fully pre-excited" QRS complexes. Furthermore, in the presence of more than one accessory connexon further possibilities exist for the generation of tachycardias with broad QRS complexes not caused by bundle branch block. These tachycardias may cause considerable diagnostic confusion and therapeutic difficulty. Several questions may be defined: (a) the mechanism of tachycardia; (b) the location and type of the accessory pathway; and (c) the functional relation of the pathway to the tachycardia circuit. In this report we describe diverse mechanisms of pre-excited tachycardia in 12 patients and how these mechanisms can be elucidated and distinguished using electrophysiological techniques and sometimes antiarrhythmic drugs with specific effects.

Patients and methods

Twelve patients aged 7–62 (median 20) years were referred for investigation of frequent (>10 a year) recurrent tachycardias associated with ventricular pre-excitation. The Table summarises the relevant details. Electrophysiological studies were performed in all patients as described previously.2 In addition to right heart sites, recordings were obtained from the left atrium or coronary sinus in all but three patients (cases 1, 7, and 8). No patient had taken any antiarrhythmic drug for 72 hours before the procedure. None of the patients had received amiodarone. The diagnosis of the cause of ventricular pre-excitation was made according to previously described criteria.3 4 Re-entry junctional tachycardia is defined

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Accepted for publication 5 June 1984

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as tachycardia due to circus movement using pathways connecting atrial and ventricular myocardium—that is, atrioventricular node-His pathway, direct atrioventricular accessory connexion.

**Results**

The Table summarises the individual results. The electrophysiological studies indicated the cause of ventricular pre-excitation in all patients (Table). During the study 21 tachycardias were induced in 12 patients. Three patients had three different tachycardias, three had two types, and six had a single mechanism. During 13 tachycardias the QRS complex was broad and resembled ventricular pre-excitation during sinus rhythm or atrial pacing. These tachycardias were considered to show ventricular pre-excitation. An accessory pathway supported pre-excited tachycardia in only six of 13 instances.

**NODOVENTRICULAR PRE-EXCITATION**

In six patients (cases 5 and 7–11) ventricular pre-excitation during tachycardia was the result of nodovenous conduction. In five of these atrioventricular nodal re-entry was the cause of tachycardia. Three patients have been described in detail in previous communications.

**ATRIOVENTRICULAR PRE-EXCITATION**

In seven patients ventricular pre-excitation during tachycardia was caused by conduction over a direct accessory pathway. Three patients had simple antidromic tachycardia with anterograde conduction over the accessory pathway and retrograde re-entrant conduction over the atrioventricular node. Verapamil 0-1 mg/kg intravenously was given during this type of tachycardia in one patient (case 4). Slowing of tachycardia as a result of an increase in the retrograde conduction time preceded termination by retrograde block after one minute. This information supports involvement of the atrioventricular node in the retro-
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Fig. 1 Case 5: electrograms and electrocardiograms recorded during the atrial extrastimulus method. (a) During regular atrial pacing at a cycle length of 620 ms the HV interval is zero. Atrial premature stimulation (S₂) results in normal prolongation of the AH interval with no change in the HV interval. (b) An earlier stimulus conducts with a sudden increase in AH interval, normal HV interval (40 ms), and normal QRS. These findings indicate the presence of dual AH pathways and a nodoventricular pathway or fasciculoventricular pathway in close association with the fast pathway. HRAE, high right atrial electrogram; DCSE, distal coronary sinus electrogram; PCSE, proximal coronary sinus electrogram; HBE, His bundle electrogram; A atrial, H His, and V ventricular (electrograms). Paper speed 100 mm/s. (See also Fig. 2.)

grade limb of tachycardia. One of these patients (case 1) had several additional forms of tachycardia. Narrow complex tachycardias of two distinct cycle lengths were observed. Different AH intervals accounted for the differing cycle lengths. On some occasions partial pre-excitation was observed during these tachycardias. Disappearance of partial pre-excitation was associated with an increase in rate due to shortening of the AH interval. Conversion of a narrow or partially pre-excited QRS complex tachycardia to a broad complex tachycardia could be achieved with single ventricular premature stimuli (Fig. 6a). The QRS morphology during this broad complex tachycardia suggested right ventricular pre-excitation. The uniformity of the tachycardia cycle length regardless of the cycle length of the preceding narrow tachycardia suggested a different mechanism (Fig. 6b). Although narrow complex tachycardias could not be terminated by atrial underdrive pacing, broad complex tachycardias were easily terminated with this method. Verapamil 0.1 mg/kg given intravenously as a bolus during the wide complex tachycardia had no effect. In contrast, flecainide (2 mg/kg) given over five minutes terminated tachycardia in the anterograde limb. These data strongly suggest the presence of a direct atrioventricular accessory connexion supporting the anterograde limb of fully pre-excited tachycardia. This pathway probably acted as a bystander during the narrow QRS tachycardias of intranodal origin. In both types of tachycardia the retrograde sequence was identical, with early low right atrial activation. Alternatively, a second concealed accessory pathway may have been the retrograde limb of all tachycardias. This would explain the total absence of any effect of verapamil, which may terminate simple antidiromic tachycardias in the retrograde limb as in the patient in case 4. Atrioventricular nodal tachycardia with incidental conduction over a direct right sided accessory pathway occurred in one other patient. This patient also had orthod-
romic atrioventricular re-entrant tachycardia (Fig. 7). Two patients (cases 3 and 6) had tachycardias which did not involve the atrioventricular node. In both patients anterograde conduction occurred over a right accessory pathway and retrograde conduction over a left concealed accessory pathway. In one patient this situation was transient. In the other (case 3) intravenous verapamil (0.1 mg/kg) had no effect on this type of tachycardia, although it terminated the typical orthodromic variety. On another occasion, intravenous flecainide readily terminated tachycardia involving two accessory pathways.

The Table shows the final treatment of each patient. One patient (case 6) underwent surgical ablation of the septal accessory atrioventricular pathway and His bundle. The right sided accessory pathway could not be ablated. Tachycardias have not recurred. In the patient in case 12 the right accessory pathway was successfully located and ablated. Three patients had implanted antitachycardia pacemakers, and in the remainder the tachycardias were controlled with antiarrhythmic drugs.

**Discussion**

The most frequent cause of wide complex tachycardia in the Wolff-Parkinson-White syndrome is atrioventricular re-entrant tachycardia with rate related bundle branch delay or block. In this analysis we have not considered these tachycardias but rather those in which the wide QRS complex during tachycardia was accounted for by conduction over an accessory pathway resulting in ventricular pre-excitation. This may occur in one of two ways: (a) the accessory connexion is part of the tachycardia circuit or (b) it is activated incidentally by tachycardia occurring as a result of some independent mechanism. The most usual cause of the phenomenon is the first, in which activation proceeds from atria to ventricles over an accessory pathway and returns via the atrioventricular node or...
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Fig. 4 Diagrams of postulated mechanisms of tachycardia in Fig. 3. (a) The initial event is a change in the relative timing of the His potential independent of a change in the ventricular cycle. The sudden advancement of the His bundle may thus have occurred after engagement of the nodoventricular pathway (NVP). A faster intranodal pathway (open arrow) distal to the accessory pathway accounts for the shorter HV interval, the earlier timing of the next atrial electrogram and the subsequent QRS complex, and the shorter tachycardia cycle length. As the His bundle is not on circuit in this mechanism changes in the HV interval are not necessarily exactly reflected by changes in cycle length. In the no stable tachycardia the AV interval has increased by 30 ms without affecting the HV relation indicating some delay in the anterograde limb proximal to the accessory connexion, which possibly accounts for the difference between VH shortening and tachycardia cycle decrease. (b) A possible mechanism of re-entrant tachycardia involves the nodoventricular pathway. The nodoventricular pathway may be in close association with the AV node (AVN). Pre-excitation during tachycardia could be due to anterograde conduction over the nodoventricular pathway. This pathway may insert into the myocardium or the right bundle branch. Retrograde conduction via the His-Purkinje system to the atrioventricular node completes a potential circuit. To explain an HV interval of zero the nodoventricular pathway must insert directly into the right bundle branch (as drawn). If more than one retrograde pathway exists (distant His-Purkinje branches) the activation of the His bundle could vary with respect to the onset of the QRS (varying HV interval). This model, however, does not explain all the features observed in the patient in case 11. RBB (LBB), right (left) bundle branch.

any other available pathway. The exact prevalence of this form of tachycardia is not known. Wellens et al were able to initiate this tachycardia in four of 87 patients undergoing electrophysiological studies.7 Benditt et al, in a review of their experience of wide complex pre-excited tachycardias, noted this mechanism in seven of 26 patients.8 In three of these patients more than one accessory pathway was demonstrated. Three of our patients had simple antidromic tachycardia (cases 1, 4, and 12), and in a further two anterograde conduction occurred over one accessory connexion and the retrograde limb of the circuit was formed by a second accessory pathway. In one patient the two pathways were capable of anterograde and retrograde conduction, a finding reported only once previously.9

Although the combination of dual AH pathways and accessory atrioventricular pathways is well recognised,10 there are few reports of intranodal re-entrant tachycardia with incidental conduction over an accessory atrioventricular connexion. Sudden onset and termination of pre-excitation without changes in tachycardia cycle length or atrial sequence are highly suggestive of incidental pre-excitation.11 Furthermore, varying degrees of pre-excitation during tachycardia exclude anterograde on-circuit activation of the accessory pathway, which must therefore be a bystander.

The role of nodoventricular pathways in tachycardias has been discussed at length in published reports.11–13 The central question is whether or not these pathways support tachycardias or act as bystanders. Gallagher et al13 have presented strong arguments that nodoventricular conduction during tachycardia indicates the involvement of this pathway in the anterograde limb of what is essentially a ventricular or subjunctional tachycardia14 incorporating accessory and intraventricular conduction pathways. In several instances they showed ventriculoatrial dissociation during tachycardia, which, although supportive of subjunctional macroreentry, does not exclude intranodal re-entry.15 The macroleernt mechanism is undoubtedly possible, but as pointed out by Akhtar6 no unequivocal proof as yet exists. Two of our patients illustrate this controversy. The findings in one patient (case 5) favour an intranodal rather than a subjunctional mechanism (Figs. 1 and 2). These findings strongly resemble those we have found in other patients.4,5 Another patient (case 11) with features of accessory nodoventricular conduction showed unusual phenomena, which could be interpreted as reflecting either a junctional or subjunctional mechanism of tachycardia. During tachycardia the relation of the His potential to the QRS (HV interval) changed suddenly (Figs. 3, 4, and 5). Kuck et al noticed similar changes during simple antidromic tachycardias which were attributable to equal changes in the VH interval.16 Trantham et al have reported three patients with findings similar to those in our patient.17 Each case showed tachycardia with either long or short VH intervals. These findings were interpreted as consistent with the hypothesis that the tachycardia circuit comprised anterogarde nodoventricular conduction and retrograde His-Purkinje conduction (Fig. 4b). They suggested that changes in intraventricular conduction time accounted for both
Fig. 5 Case 10: electrograms and electrocardiograms showing the initiation of tachycardia. During regular atrial pacing the QRS complexes show left bundle branch block morphology and a short HV interval. An atrial extrasystole conducts with a normal HV interval (35 ms) and a normal QRS complex, and tachycardia is initiated. Earliest atrial activation during tachycardia is seen on the coronary sinus electrogram (PCSE). After several complexes the HV interval shortens because pre-excitation returns. These findings indicate anterograde atrioventricular nodal and retrograde direct accessory ventriculoatrial conduction sustaining tachycardia with incidental nodoventricular conduction independent of the tachycardia mechanism. Abbreviations as in Fig. 1.

Fig. 6 (a) Case 1; electrograms and electrocardiograms during stable tachycardia. Initially tachycardia shows partial pre-excitation with an HV interval of 25 ms and cycle length alternation of 290/320 ms. A single right ventricular stimulus results in conversion to tachycardia with fully pre-excited QRS complexes and a stable cycle length of 310 ms. This phenomenon was also observed during slower narrow complex tachycardia. The cycle length of the resulting wide complex tachycardia was always 300–310 ms regardless of the initial cycle length. (b) Diagram showing changes in tachycardia cycle length produced by a single ventricular stimulus. The narrow or partially pre-excited tachycardias fell into slow and fast categories. These tachycardias showed varying degrees of pre-excitation which on occasion could be related to changes in the AH interval. This group of tachycardias was thought to reflect atrioventricular nodal re-entry with variable degrees of incidental pre-excitation over a right accessory pathway. Conversion to wide complex tachycardia was associated with uniformity of cycle length regardless of the initial tachycardia. The retrograde sequence remains unaltered. The wide complex tachycardia was considered to reflect atrioventricular re-entry with anterograde conduction over the right accessory pathway. The retrograde limb is either the normal atrioventricular node or an additional accessory pathway. It is suggested that the ventricular premature stimulus responsible for the conversion conducted retrogradely with consequent earlier anterograde accessory pathway activation and incorporation in the circuit.
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The change in the VH interval and the tachycardia cycle length. It is of interest to note, however, that the change in VH interval in each case exceeds the change in tachycardia cycle length just as in our patient. The increase in cycle length should equal that of the VH interval unless there is consequent delay in another part of the circuit such as the atrioventricular node. Thus shortening of the VH interval may be followed by increasing intranodal delay resulting in a shortening of tachycardia cycle length somewhat less than the change in the VH interval. This delay is most likely to occur in that part of the postulated circuit between the His bundle and the nodoventricular pathway. If this were so, however, a longer HA interval would be expected during faster stable tachycardia reflecting additional retrograde intranodal delay. In our patient this was not seen. To account for the VH interval of zero during tachycardia it would be necessary to postulate insertion of the nodoventricular fibre into the right bundle branch with anterograde conduction time to the myocardium equal to retrograde conduction time to the His bundle. Such a mechanism would also be expected to operate during atrial rhythms if anterograde His activation were sufficiently delayed relative to ventricular activation. Since negative HV values were seen during atrial extrastimulation, this mechanism need not be invoked in this particular instance. Nodoventricular pathways may have complex relations with intranodal structures. For example, such pathways appear to be uniquely functionally associated with one of two intranodal pathways as observed in one of our patients (case 5). The observations in the patient in case 12 may reflect a more complex interrelation between accessory and normal pathways. Morady et al described a very similar patient who appeared to have a nodoventricular pathway in association with a slow intranodal pathway. In their case, there was HV dissociation during pre-excited tachycardia excluding intraventricular macroreentry. They argued that tachycardia reflected intranodal re-entry.

The combination of incidental nodoventricular conduction during atrioventricular re-entrant tachycardia involving a direct accessory pathway as in the patient in case 11 has also been described by Motte et al and later by Lerman et al. In the latter report, the authors propose a macroreentry circuit comprising the nodoventricular pathway in the anterograde limb and a septal concealed accessory pathway in the retrograde limb. Nevertheless, the HV
interval during tachycardia is 25 ms suggesting anterograde activation of the His bundle and participation of both the normal and accessory pathways within the circuit with anterograde conduction in parallel over both pathways. The presence of varying fusion complexes at the onset of tachycardia without any associated changes in cycle length implies that the nodoventricular pathway is not on circuit.

Seven of the patients in this series had more than one type of tachycardia. In four patients at least two distinct different mechanisms could be identified. In two, possibly three, patients a second accessory pathway was present. Multiple accessory pathways, although uncommon, are a well recognised cause of complex junctional re-entrant mechanisms.\(^9\)\(^{24}\) Their presence is, however, often difficult to establish. Involvement of two such pathways in a tachycardia circuit excluding the atrioventricular node is rare,\(^7\)\(^{9}\)\(^{25}\) and if verapamil fails to affect tachycardia an erroneous diagnosis of ventricular tachycardia may be made.

In conclusion, broad complex tachycardias associated with ventricular pre-excitation may be caused by a multitude of different mechanisms and are often associated with complex junctional substrates—that is, more than one abnormal pathway. Pre-excitation during junctional tachycardia does not imply participation of an accessory pathway in the circuit. Several varieties of tachycardia mechanism may coexist and interact with each other. These arrhythmias may be misdiagnosed as ventricular tachycardia with consequent inappropriate investigations. The use of antiarrhythmic drugs with known effects (for example, verapamil, flecaïnid) is valuable in assessing these complex arrhythmias and should be considered during the electrophysiological study. Accurate diagnosis of the various diverse mechanisms is not possible without detailed studies, which may provide essential information on which to base treatment, whether it is medical, electrical, or surgical, or in particular, uses transvenous techniques designed to ablate accessory conduction.\(^{26}\) With regard to the last two approaches, determination of the role of the accessory pathway is essential.

We thank Dr A W Nathan for supplying details of the patient in case 10.

References


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D E Ward, D H Bennett and J Camm

Br Heart J 1984 52: 369-376
doi: 10.1136/hrt.52.4.369

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