Selective blockade of retrograde fast pathway by intravenous disopyramide in paroxysmal supraventricular tachycardia mediated by dual atrioventricular nodal pathways

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SUMMARY Electrophysiological effects of 2 to 2.5 mg/kg iv disopyramide were studied in 10 patients with dual nodal pathways who used a slow pathway for anterograde and a fast pathway for retrograde conduction during paroxysmal supraventricular tachycardia (mean cycle length 308.5±37 ms; range 260–370 ms). Disopyramide terminated the tachycardia in six cases by production of ventriculoatrial block in five and by sinus overdrive in one. In the remaining four patients cycle length of the paroxysmal supraventricular tachycardia increased significantly from 270±8 ms to 377.5±28 ms. In all 10 patients disopyramide depressed retrograde fast pathway conduction manifest by an increase in mean ventricular paced cycle length producing ventriculoatrial block from ≤296.5±25 ms to 358±60 ms, and increase in retrograde fast pathway effective refractory period from ≤246±34 ms to 325±36 ms; the drug abolished ventriculoatrial conduction in two cases.

Anterograde slow pathway and fast pathway conduction properties were unchanged after disopyramide (atrial paced cycle length producing AH block 292±30 to 306.5±30 ms; effective refractory period of anterograde fast pathway ≤274±56 to 284±44 ms, before and after the drug, respectively) suggesting that anterograde conduction was not crucial either for sustainment or for failure to initiate paroxysmal supraventricular tachycardia after the drug.

Paroxysmal supraventricular tachycardia could not be reinduced in six cases after disopyramide. In the other four the ventricular paced cycle lengths producing ventriculoatrial block (318±41 ms) and effective refractory period of retrograde fast pathway (320±28 ms) were shorter than the cycle length of reinduced paroxysmal supraventricular tachycardia (367.5±35 ms) allowing perpetuation of the tachycardia.

We conclude that disopyramide breaks atrioventricular nodal re-entrant tachycardia by specific blockade of the retrograde fast pathway though the effect on anterograde atrioventricular nodal conduction is variable.

Electrophysiological studies have shown that longitudinal dissociation of the atrioventricular node into dual pathways constitutes the most common mechanism of re-entry in the mediation of paroxysmal supraventricular tachycardia. Usually the tachycardia uses the slow pathway for anterograde conduction and the fast pathway for the return impulse; and the three most commonly recommended drugs, that is verapamil, digitalis, and propranolol, inhibit the arrhythmia by depression of conduction through the anterograde slow pathway. Some studies have also reported the effects of procainamide and quinidine on laboratory induction of paroxysmal supraventricular tachycardia.

Disopyramide phosphate, a synthetic compound with antiarrhythmic actions similar to those of quinidine, was introduced by Mokler and Van Arman in 1962. Because of its demonstrably variable effects on the atrioventricular nodal conduction and refractory periods it was not construed to be effective in paroxysmal supraventricular tachycardia mediated by dissociation of the atrioventricular node into dual...
**Disopyramide in paroxysmal supraventricular tachycardia**

pathways.\(^{10}\)\(^ {11}\) On the other hand, in patients with paroxysmal supraventricular tachycardia associated with Wolff-Parkinson-White syndrome, disopyramide inhibits both anterograde and retrograde conduction through the bypass tract\(^ {10}\)\(^ {12}\) and has been recommended for use in paroxysmal supraventricular tachycardia using atioventricular node bypass tract. A few studies conducted in a small cohort of patients without overt pre-excitation have shown that intravenous disopyramide terminates 33 to 68% of acute attacks of paroxysmal supraventricular tachycardia.\(^ {13}\)\(^ {16}\) No intracardiac electrophysiological studies to delineate the mechanism of paroxysmal supraventricular tachycardia or its mode of termination were, however, performed in any of these investigations. It is well recognised that concealed atioventricular accessory pathways, functionally silent during sinus rhythm and therefore clinically unsuspected, participate in 15 to 30% cases of re-entrant paroxysmal supraventricular tachycardia\(^ {17}\); hence it becomes obligatory to rule out participation of this circuit in evaluating the effect of drugs on atioventricular nodal re-entrant paroxysmal supraventricular tachycardia. The efficacy of intravenous disopyramide in terminating episodes of paroxysmal supraventricular tachycardia in a select group of patients with dual atioventricular nodal pathway mediation has thus far not been adequately investigated, and formed the aim of the present study.

**Subjects and methods**

Ten patients, four men and six women, aged 25 to 56 years (mean 42.8 years) comprised the study material selected on the following criteria: (1) a history of electrocardiographic documented recurrent paroxysmal supraventricular tachycardia, (2) absence of pre-excitation during sinus rhythm in all available electrocardiograms, (3) electrophysiological documentation of longitudinal dissociation of the atioventricular node during paroxysmal supraventricular tachycardia, and (4) exclusion of a retrogradely active concealed bypass tract during paroxysmal supraventricular tachycardia by intracardiac catheter techniques.

Electrophysiological studies were performed in the post-absorptive non-sedated state a week after discontinuation of all cardioactive medications. Informed written consent was obtained from all. One patient had angina of effort; others were free of organic heart disease.

Six 6F bipolar catheters were introduced transvenously and positioned at high, mid, and low right atrium, coronary sinus, tricuspid valve (for His bundle electrogram), and right ventricular apex. The mid-right atrial and right ventricular electrodes were used for pacing and programmed stimulation. The electrograms from the high and low right atrium, tricuspid valve, and coronary sinus were recorded (frequency range 30 to 500 Hz) together with multiple scalar leads (combinations of lead I, II, III, aVF, or V1) on VR-12 photographic recorder (Electronics for Medicine) or on Mingograf 8 channel ink jet recorder (Siemens-Elema) at paper speeds of 100 and 250 mm/s. Intracardiac stimulation was performed at approximately twice the diastolic threshold by a battery powered stimulator (Digitimer arrhythmia investigating system, Neurolog-4279) producing square wave pulses of 2 ms duration.

The study consisted of evaluation of anterograde and retrograde conduction properties, refractory periods of the slow and fast pathways, and the mechanism of paroxysmal supraventricular tachycardia, pursued in the following sequence: (a) incremental atrial pacing to a paced cycle length producing atrioventricular block; (b) programmed atrial extrastimulus testing at decreasing coupling intervals using single \(A_1 A_2\) extrastimuli during sinus rhythm and during one or more atrial paced cycle lengths. Double extrastimuli \(A_1 A_2 A_3\) were given when single stimuli failed to induce paroxysmal supraventricular tachycardia; (c) incremental ventricular pacing to a paced cycle length producing ventriculoatrial block; (d) programmed ventricular extrastimulation at decremental coupling intervals at a driven ventricular cycle length just shorter than sinus cycle length; (e) induction of paroxysmal supraventricular tachycardia by rapid atrial pacing, atrial or ventricular extrastimulation, and delineation of the limbs of the re-entrant circuit by analysis of the mode of initiation of paroxysmal supraventricular tachycardia, anterograde and retrograde conduction times, and sequence of retrograde atrial activation (from endocardial map recorded from multiple atrial sites); and (f) ventricular programmed extrastimulation during induced paroxysmal supraventricular tachycardia to rule out concealed septal bypass tract participation.

After the verification of dual atioventricular nodal pathway participation during paroxysmal supraventricular tachycardia, 2 mg/kg of disopyramide phosphate was injected intravenously over a period of three minutes. Cuff blood pressure was closely monitored. Intracardiac and surface electrograms were continuously recorded at paper speed of 50 mm/s from the beginning of injection in order to study the mode of termination of the tachycardia. If the tachycardia failed to break within 10 minutes after initiation of injection, and if no side effects were encountered, an additional dose of 0.5 mg/kg was administered intravenously over one minute (four patients). If the paroxysmal supraventricular tachycardia was not abolished in the following five minutes, it was terminated electrically by atrial extr-
stimulus or atrial overdrive pacing.
After conversion to sinus rhythm by either mode, the cycle length, AH, and HV intervals were recorded at paper speed of 100 mm/s. Atrial and ventricular pacing and extrastimulation studies were then conducted in the sequence described earlier and an attempt was made to reinduce paroxysmal supraventricular tachycardia. The post-drug studies were completed within 30 minutes of termination of the arrhythmia.

ELECTROPHYSIOLOGICAL DEFINITIONS
RA1, CS1, A1, H1, and V1 are high right atrial, left atrial (recorded from coronary sinus), low septal right atrial, His bundle, and ventricular responses during sinus rhythm or driven atrial or ventricular stimuli (S1). RA2, CS2, A2, H2, and V2 are respective responses to the extrastimulus (S2). Conduction intervals and refractory periods were measured as defined by Wu et al. Anterograde and retrograde block rates were defined as the longest paced cycle length (atrial and ventricular, respectively) which failed to conduct to the distal chamber. In an occasional case rapid pacing carried out to very high rates did not achieve block. In such cases the highest paced rate tested was considered to be the "block rate" for purposes of analysis. This occurred only during the control studies and not after the drug so that the magnitude of decrease in block rates attributed to disopyramide in these cases indicates the minimal values of decrement.

Diagnosis of atrioventricular nodal re-entrant paroxysmal supraventricular tachycardia was made using combinations of the following criteria.
(1) Induction of tachycardia related to achievement of a critical AH delay, with both incremental atrial pacing and atrial extrastimulation (10 patients).\(^1\)\(^{-21}\)
(2) Demonstration of discontinuous A1A2 to H1H2 curves, or two different AH intervals at identical pacing cycle length, suggestive of dual atrioventricular nodal pathways with induction of tachycardia related to anterograde block in the fast pathway (seven patients).\(^1\)\(^{-21}\)
Because it is possible to fulfil criteria (1) and (2) in patients with concealed extranodal bypass tracts, one or more of the following additional criteria were sought for the diagnosis of atrioventricular nodal re-entry.
(3) Normal retrograde atrial activation sequence during tachycardia, namely, low septal right atrium activated earlier than all other atrial sites (10 patients).\(^2\)
(4) Demonstration of atrial activation simultaneous with onset of ventricular activation during tachycardia, suggesting that the ventricle is not an essential component of the re-entry circuit (10 patients).\(^2\)
(5) Increase in ventriculoatrial interval with incremental ventricular pacing producing Mobitz type I block at a critical rate, suggesting retrograde atrioventricular nodal conduction (9 patients).\(^1\)\(^{-2}\)
(6) Demonstration of His bundle activation (H2) preceding atrial activation (A2) with ventricular extrastimulus or pacing, suggesting retrograde atrioventricular nodal conduction (four patients).\(^2\)
(7) Exclusion of concealed septal bypass tract by showing failure of programmed ventricular extrastimulation to pre-excite the atrium when delivered at the critical time when the His bundle was rendered refractory by anterograde depolarisation during the normal course of paroxysmal supraventricular tachycardia.

Patients with concealed bypass tract participation during the tachycardia\(^2\) were excluded from the study.

Results
The results are set out in the Table. The mean basal cycle length during sinus rhythm was 578±107 ms (range 440 to 740 ms); the mean AH interval was 78.8±13 ms (range 60 to 90 ms) and the mean HV time was 44±6 ms (range 35 to 50 ms).

INDUCTION OF PAROXYSMAL SUPRAVENTRICULAR TACHYCARDIA
Sustained paroxysmal supraventricular tachycardia could be induced in all 10 patients: by rapid atrial pacing in seven, by single atrial extrastimulus in three, by double atrial extrastimulus in two, and by rapid ventricular pacing in one case. Induction of paroxysmal supraventricular tachycardia by atrial stimulation was related to achievement of a critical AH prolongation consequent to anterograde block of the fast pathway and the resultant atrioventricular conduction via the slow pathway. Induction of paroxysmal supraventricular tachycardia by ventricular pacing was obtained without a ventriculoatrial delay, reflecting block in the retrograde slow pathway.

Cycle length of the tachycardia ranged between 260 and 370 ms (mean 308.5±37 ms). All patients showed anterograde conduction through the slow pathway and retrograde conduction through the fast pathway.

TERMINATION OF PAROXYSMAL SUPRAVENTRICULAR TACHYCARDIA
Disopyramide could terminate paroxysmal supraventricular tachycardia in six cases. The break occurred immediately on completion of injection in three cases, and between one minute 30 seconds and four minutes
Disopyramide in paroxysmal supraventricular tachycardia

Effect of disopyramide on paroxysmal supraventricular tachycardia

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VT, paroxysmal supraventricular tachycardia; CL, cycle length; ERP, effective refractory period; FP-ERP, effective refractory period of the pathway; CL-AVB, longest atrial paced cycle length producing atrioventricular block; CL-VAB, longest ventricular paced cycle length producing ventriculoatrial block; C, control; D, after intravenous disopyramide.

48 seconds (mean two minutes six seconds) after the injection in the other three. The break was achieved by ventriculoatrial block (retrograde fast pathway block) in five patients (cases 1, 2, 3, 4, and 10) (Fig. 1) and by sinus overdrive (described below) in the sixth case. Among the five cases showing ventriculoatrial block, the cycle length of paroxysmal supraventricular tachycardia before termination remained unchanged in three (cases 1, 4, and 10) and increased from 315 ms (mean) to 390 ms (mean) in two (cases 2 and 3).

In the sixth patient (case 5) the arrhythmia was converted in a peculiar manner. The drug not only increased the cycle length of the tachycardia (from 340 to 380 ms) but also accelerated the sinus cycle length to 370 ms, producing an isorhythmic "sinus-paroxysmal supraventricular tachycardia dissociation" as evidenced by activation of low septal right atrium by the retrograde impulse of the tachycardia and stimulation of the high right atrium, left atrium, and an unknown area of atrial myocardium by sinus impulse until such time that the latter could penetrate the atrioventricular node and overdrive the paroxysmal supraventricular tachycardia (Fig. 2).

Conversion of the tachycardia was succeeded by sinus rhythm in four cases (cases 3, 4, 5, and 10). One patient (case 1) showed irregular sinus cycles with QRS aberrancy with junctional escapes followed by a sudden change to regular sinus rhythm, and another (case 2) developed junctional escape succeeded by sinus capture and sinus rhythm.

In four patients (cases 6, 7, 8, and 9) paroxysmal supraventricular tachycardia could not be abolished despite administration of the additional dose of the

Fig. 1 Case 1. Termination of paroxysmal supraventricular tachycardia by abrupt ventriculoatrial block. The cycle length of paroxysmal supraventricular tachycardia is constant (320 ms) until the seventh cycle where "V" is not followed by "A" (arrow). The first sinus beat shows aberrant conduction. Abbreviations as in text.
The cycle length of the tachycardia, however, increased significantly from 270±8 ms to 377-5±28 ms (p<0.01).

Cycle length during sinus rhythm immediately after termination of paroxysmal supraventricular tachycardia was 509-5±66 ms (range 395 to 640 ms), not significantly different from the basal sinus cycle length of 578±107 ms (NS). The AH interval of 84-5±14 ms (range 60 to 105 ms) after the drug was also not significantly different (NS). The HV interval was, however, significantly prolonged to 54-5±10-6 ms (range 40-70 ms) (p<0.01). This increase was most pronounced in cases receiving the additional dose of the drug (Table).

**Fig. 2** Case 5. Termination of paroxysmal supraventricular tachycardia by sinus overdrive. Top panel: paper speed 50 mm/s; left, five beats of paroxysmal supraventricular tachycardia (cycle length 330 ms) before disopyramide. The atrial activation in the coronary sinus electrogram immediately follows ventricular activation. Right, after administration of the drug, the “A” and “V” complexes gradually get superimposed on each other, suggesting isorhythmic atrioventricular dissociation.

Bottom panel: paper speed 100 mm/s: in a later part of the continuous record the paroxysmal supraventricular tachycardia cycle length varies from 330 to 380 ms, while the atrial cycle length is constant at 370 to 380 ms. The atrial deflection at the high right atrium (HRA) precedes that at the His bundle electrogram and coronary sinus, indicating its origin in the sinus node. “Sinus to paroxysmal supraventricular tachycardia dissociation” is clearly seen, till sinus impulse captures the ventricle with an AH interval of 130 ms (arrow) converting the paroxysmal supraventricular tachycardia into sinus rhythm (cycle length 380 ms, AH 90 ms, HV 60 ms).

**Anterograde Conduction Properties**

The effective refractory period of the atrium was 234±32 ms (range 190 to 280 ms) before, and 250±39 ms (range 180 to 310 ms) after the drug (NS). The atrial paced cycle length producing AH Wenckebach block was 292±30 ms (range 240 to 330 ms) before, and 306-5±30 ms (range 250 to 340 ms) after disopyramide (NS). The atrial paced cycle length producing atrioventricular block increased in seven, decreased in one (case 5), and remained unchanged in two patients (Fig. 3). The cycle length of paroxysmal supraventricular tachycardia (308-5±37 ms) was longer than the paced cycle length that produced atrioventricular nodal block (292±30 ms) during the
Disopyramide in paroxysmal supraventricular tachycardia

Fig. 3 Effect of disopyramide on atrial paced cycle length producing atrioventricular block.

Control study.

The effective refractory period of the anterograde fast pathway was \( \leq 274 \pm 56 \text{ ms} \) (range \( <200 \) to \( 400 \) ms) before, and \( \leq 284 \pm 44 \text{ ms} \) (range \( <220 \) to \( 360 \) ms) after the drug (NS) (Fig. 4). The effective refractory period of the anterograde slow pathway could not be compared before and after disopyramide because atrioventricular nodal conduction was limited by atrial refractoriness before and/or after the drug.

RETROGRADE CONDUCTION PROPERTIES

Disopyramide increased the effective refractory period of the ventricle from \( 215 \pm 12 \text{ ms} \) (range \( 200 \) to \( 230 \) ms) to \( 229 \pm 11 \text{ ms} \) (range \( 210 \) to \( 240 \) ms) \((p<0.05)\). During the control study the ventricular paced cycle length that produced ventriculoatrial block ranged from \( <260 \) to \( 325 \text{ ms} \) (mean \( \sim 296 \pm 5 \pm 25 \text{ ms} \)). After disopyramide the ventricular paced cycle length that produced ventriculoatrial block increased in all the patients irrespective of whether or not the paroxysmal supraventricular tachycardia was terminated by the drug (mean \( 358 \pm 60 \text{ ms} \); range \( 286 \) to \( 470 \) ms) \((p<0.01)\) (Fig. 5). In two cases (1 and 4) ventricular pacing at a cycle length just shorter than the sinus cycle length disclosed complete abolition of ventriculoatrial conduction (Fig. 6). The ventriculoatrial conduction times at identical pacing rates were longer after disopyramide compared with control in all the cases. In all the subjects the cycle length of paroxysmal supraventricular tachycardia \( (308 \pm 37 \text{ ms}) \) was longer than the cycle length that produced ventriculoatrial block before disopyramide \( (296 \pm 25 \text{ ms}) \).

The effective refractory period of the retrograde fast pathway before disopyramide was \( \sim 246 \pm 34 \text{ ms} \) (range \( <210 \) to \( 300 \) ms). After disopyramide the effec-

Fig. 4 Effect of disopyramide on the effective refractory period of anterograde (left panel) and retrograde (right panel) fast pathway. The horizontal interrupted lines represent two cases in which ventriculoatrial conduction was abolished by the drug.

Fig. 5 Effect of disopyramide on ventricular paced cycle length producing ventriculoatrial block.
REINDUCTION OF PAROXYSMAL SUPRAVENTRICULAR TACHYCARDIA

While six subjects lost the ability to reinitiate the tachycardia, paroxysmal supraventricular tachycardia could still be induced in four (cases 5, 6, 7, and 9) after disopyramide (Fig. 7). Among the latter, three (cases 6, 7, and 9) had not converted to sinus rhythm with the drug and the fourth (case 5) had responded with sinus overdrive. In these four cases the paroxysmal supraventricular tachycardia cycle length was 290±29 ms (range 270 to 340 ms) before and 367.5±35 ms (range 330 to 410 ms) after disopyramide (p<0.05) (Fig. 8). The ventricular paced cycle length producing ventriculoatrial block increased in all four, from a mean of 285±26 ms to 318±41 ms after the drug (Fig. 9). Among the former six cases, paroxysmal supraventricular tachycardia was not inducible despite achieving AH intervals equal to or longer than the critical AH time that had initiated paroxysmal supraventricular tachycardia in the control state. This finding suggests that an increase in the retrograde fast pathway refractory period was responsible for failure to initiate paroxysmal supraventricular tachycardia after the drug.

During the control study, the cycle length of paroxysmal supraventricular tachycardia (308.5±37 ms) was longer than the atrial paced cycle length that produced AH block (292±30 ms) in all the patients. Likewise, in the four patients in whom sustained paroxysmal supraventricular tachycardia could be
Disopyramide in paroxysmal supraventricular tachycardia

Control

Disopyramide

Fig. 7 Case 5. (A) Before disopyramide; induction of paroxysmal supraventricular tachycardia by single atrial extrastimulus \( S_1S_2 = 210 \text{ ms} \). \( A_2 \) is conducted to the ventricle with \( AH_2 \) of 490 ms, and initiates the tachycardia. Within a few seconds the tachycardia spontaneously accelerated to a cycle length of 330 ms as shown in Fig. 2. (B) After disopyramide; driving cycle length is same as in (A). Single atrial extrastimulus \( S_1 \) delivered at a coupling interval of 240 ms conducts with \( AH_2 \) of 350 ms to initiate the paroxysmal supraventricular tachycardia. No acceleration of the paroxysmal supraventricular tachycardia was seen.

Fig. 8 Effect of disopyramide on reinduction of paroxysmal supraventricular tachycardia. Open circles represent cycle length of the tachycardia before disopyramide, and solid circles the cycle length of paroxysmal supraventricular tachycardia induced after administration of drug. Interrupted lines show cases in which paroxysmal supraventricular tachycardia could not be reinduced. Reinduced after disopyramide, the cycle length of paroxysmal supraventricular tachycardia \((367.5\pm35 \text{ ms})\) was longer than the atrial paced cycle length which produced atrioventricular block \((297.5\pm19 \text{ ms})\). Three of the six patients in whom we failed to reinduce paroxysmal supraventricular tachycardia after the drug had a cycle length of paroxysmal supraventricular tachycardia before the drug longer than the atrial paced cycle length producing AH block after the drug. These findings are consistent with the observation that the anterograde limb was not the limiting factor in the induction or sustainment of paroxysmal supraventricular tachycardia after disopyramide.

In all patients, the cycle length of paroxysmal supraventricular tachycardia \((308.5\pm37 \text{ ms})\) was longer than the ventricular paced cycle length producing ventriculoatrial block \((\leq296.5\pm25 \text{ ms})\) before disopyramide. In those six patients in whom paroxysmal supraventricular tachycardia could not be induced...
after the drug the ventricular paced cycle length that produced ventriculoatrial block after disopyramide (385±53 ms) was longer than the cycle length of paroxysmal supraventricular tachycardia (321±33 ms) during control study. In the other four patients with inducible paroxysmal supraventricular tachycardia after the drug the cycle lengths producing ventriculoatrial block (318±41 ms) and effective refractory period of the retrograde fast pathway (320±28 ms) were shorter than the post-drug cycle lengths of paroxysmal supraventricular tachycardia (367.5±35 ms), indicating that the fast pathway was not sufficiently inhibited to prevent perpetuation of paroxysmal supraventricular tachycardia.

Discussion

The initiation and sustainment of re-entrant tachycardia require the presence of a closed electrical circuit, a unidirectional block in one of the limbs, (usually manifest as a discrepancy in anterograde refractory periods of the two limbs), and a slow velocity of conduction in a part of the circuit. In patients with paroxysmal supraventricular tachycardia caused by atrioventricular nodal re-entry (longitudinal dissociation of atrioventricular node), the re-entrant circuit consists of a slow anterograde pathway and a retrograde fast (atrioventricular nodal) pathway along with a proximal and a distal common pathway (within the atrioventricular node) which conduct the re-entrant impulses. Drugs that change the refractoriness or conduction velocity of a component of the re-entrant circuit have been shown to disturb this finely adjusted echo zone and terminate the tachycardia. Such drugs have also proved useful in preventing recurrences of tachycardia during follow-up. While some reports of the beneficial effects of intravenous disopyramide in terminating acute attacks of paroxysmal supraventricular tachycardia have been published, the mechanism of termination

Fig. 9 Case 5. Same patient as in Fig. 2 and 7. While producing facilitation of anterograde conduction disopyramide depressed the retrograde conduction. The ventricular paced cycle length producing ventriculoatrial block increased from 320 ms (A) to 375 ms (B). Magnitude of depression, however, was insufficient to prevent reinduction of paroxysmal supraventricular tachycardia. A, retrograde atrial activation by ventricular paced impulse; As, atrial activation by sinus impulse.
Disopyramide in paroxysmal supraventricular tachycardia

and overall efficacy of the drug in relation to conventional and other newer antiarrhythmic drugs in a select group of patients mediating the tachycardia via dual atrioventricular nodal pathways have not so far been studied in detail. Disopyramide could convert 60% of such attacks of paroxysmal supraventricular tachycardia into normal sinus rhythm in our study material. Swiryn et al. reported that oral disopyramide could prevent induction of sustained paroxysmal supraventricular tachycardia in six out of nine cases (67%) of atrioventricular nodal re-entrance. Intravenous verapamil converts 80 to 100% of episodes of paroxysmal supraventricular tachycardia into sinus rhythm but is much less effective in preventing recurrences when administered orally3,27,28 because of considerable first pass hepatic metabolism. Other drugs that have been shown to be effective in preventing induction of atrioventricular nodal re-entrant paroxysmal supraventricular tachycardia include (percentages in parentheses): intravenous ouabain (44 to 54%),4,7 intravenous propranolol (41 to 44%),5,7 intravenous ouabain plus propranolol (58%),7 intravenous propranolol (65%),6 and oral quinidine (78%).7,8 It appears from this review of published reports that, though somewhat less effective than verapamil in converting acute attacks, intravenous disopyramide compares favourably with other drugs in controlling paroxysmal supraventricular tachycardia mediated by dual atrioventricular nodal pathways and may possess the additional advantage of effectively preventing recurrences on chronic oral treatment.26 We have not attempted specifically to test in our study whether chronic oral administration of disopyramide prevents recurrence of paroxysmal supraventricular tachycardia, but such proof in a large number of patients with dissociation of the atrioventricular node into dual pathways and paroxysmal supraventricular tachycardia is desirable.

As observed in our study, Befeler et al.29 have also reported no change or shortening of the atrial effective and functional refractory periods after intravenous administration of disopyramide. On the other hand, other investigators10,30,31 found increases in these variables. An increase in ventricular effective refractory period has also been shown,10 but antegrade atrioventricular conduction has been shown to be depressed, unaffected, or even facilitated by the drug.10,29,30,32 Both antegrade slow pathway conduction (atrial paced cycle length producing atrioventricular block) and effective refractory period of antegrade fast pathway were not significantly affected in the present study.

In sharp contrast to its effect on the antegrade conduction through the atrioventricular node, disopyramide selectively and uniformly depressed conduction through the retrograde fast pathway in all the patients, as shown by the lowering of ventricular paced rate producing ventriculoatrial block, and an increase in the effective refractory period of the retrograde fast pathway. This effect was responsible for termination of paroxysmal supraventricular tachycardia in five patients (ventriculoatrial block) and prevention of reinduction of paroxysmal supraventricular tachycardia in six cases as shown by failure to induce atrioventricular nodal re-entrant echo despite achievement of an AH interval equivalent to or longer than the one that precipitated paroxysmal supraventricular tachycardia in the control state. It is clear from our data that the fast pathway, while unaffected in the anterograde direction, was rendered incapable of conducting retrogradely. Though the nature of this retrograde pathway is subject to some discussion, the available evidence indicates that it is located intranodally and is identical to the anterograde fast pathway.33

The results of other studies on the effect of oral quinidine,7,8 and oral disopyramide26 on induction of atrioventricular nodal re-entrant tachycardia corroborate the data obtained by us on intravenous disopyramide on electrophysiologically confirmed dissociation of atrioventricular nodal pathways mediated paroxysmal supraventricular tachycardia. In our study disopyramide did marginally depress the anterograde slow pathway conduction in some patients but this did not appear to be a limiting factor in inducing or sustaining the tachycardia. This is in contradistinction to the actions of intravenous verapamil,3,27,28 digitalis,4,7 and propranolol5,7 which, by and large, increase the anterograde slow pathway refractoriness and only occasionally depress the retrograde fast pathway conduction. Intravenous procainamide prevents induction of paroxysmal supraventricular tachycardia in some patients6 by inhibiting the retrograde fast pathway but may have the disadvantage of decreasing the anterograde slow pathway refractoriness (vagolytic effect) in some patients. Atrioventricular conduction was facilitated by intravenous disopyramide in one of our patients (case 5) who overdrove the paroxysmal supraventricular tachycardia by sinus tachycardia. Alterations in the sinus rate after disopyramide therapy are variable and related to a complex interaction between the direct depressant action on sinus node automaticity,32 its anticholinergic effects,34 and reflex baroreceptor mediated sympathomimetic effect caused by peripheral vasodilatation. Antegrade conduction was significantly depressed by oral disopyramide in the study of Swiryn et al.26 The discrepancy between their data and our results could possibly be the result of more prominent anticholinergic effects of the drug after intravenous administration.

Whenever disopyramide failed to terminate parox-
ysmal supraventricular tachycardia, it produced a reduction in tachycardia rate. In those four patients in whom paroxysmal supraventricular tachycardia was still inducible after disopyramide, the depression in retrograde conduction was insufficient to prevent continuation of the tachycardia. Anterograde conduction was depressed in three and facilitated in one of these cases. The cycle length of the reinduced paroxysmal supraventricular tachycardia was, however, significantly increased in all the patients.

Our data indicate that intravenous disopyramide depresses retrograde fast pathway conduction and thus terminates, or prevents induction of, paroxysmal supraventricular tachycardia in patients with atrioventricular nodal re-entrant tachycardia. Care needs to be exercised in prescribing the drug routinely in all non-specific supraventricular tachycardias because of the possibility of an increase in ventricular rate by development of 1:1 conduction through the atrioventricular node in cases of atrial flutter or paroxysmal supraventricular tachycardia caused by an ectopic atrial focus or intra-atrial re-entry in the absence of concomitant administration of atrioventricular nodal depressants (digoxin, beta blockers). This risk of 1:1 conduction on the development of atrial flutter in patients receiving disopyramide for long term prophylaxis against paroxysmal supraventricular tachycardia can be assessed during intracardiac electrophysiological study. It could not be assessed in the absence of such a test were the drug to be given empirically to all patients with intranodal paroxysmal supraventricular tachycardia because the infra-Hisian delay induced by the drug may or may not protect the ventricles from rapid response.

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Selective blockade of retrograde fast pathway by intravenous disopyramide in paroxysmal supraventricular tachycardia mediated by dual atrioventricular nodal pathways.

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