Torsades de Pointes: Electrophysiologic Studies in Patients Without Transient Pharmacologic or Metabolic Abnormalities

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SUMMARY Electrophysiologic studies were performed in 21 patients who had torsades de pointes. This ventricular tachyarrhythmia, characterized by rapid (200–250 beats/min) and irregular paroxysms and progressively varying QRS amplitude and polarity, occurred in the absence of electrolyte disturbance, antiarrhythmic drug therapy or acute ischemia. The QTc interval was prolonged in seven of 21 patients. Electrophysiologic study included ventricular pacing with the introduction of one to three extrastimuli and rapid ventricular pacing. The effect of i.v. procainamide or quinidine in these patients was also studied. Torsades de pointes was inducible in 19 of 21 patients. Induced episodes closely resembled spontaneous episodes. Torsades de pointes spontaneously progressed to ventricular tachycardia with a uniform morphology in three patients and to ventricular fibrillation in four. In eight patients, procainamide or quinidine converted torsades de pointes into typical reentrant ventricular tachycardia. Our data suggest that torsades de pointes in this setting may be a rapid reentrant ventricular tachycardia closely related to recurrent sustained ventricular tachycardia and a precursor to ventricular fibrillation and sudden death.

IN 1966, Dessertenne applied the term torsades de pointes to a distinctive ventricular tachyarrhythmia. Examples of this arrhythmia had been published under a variety of names over the preceding 50 years. It is characterized by paroxysms of ventricular tachycardia at rates typically greater than 200 beats/min in which the QRS morphology shows alternating polarity in an undulating pattern so that the complexes appear to be twisting about the baseline; hence the name torsades de pointes — “twisting of the points” (fig. 1). The designation of this arrhythmia by a specific title suggests that it is a specific electrophysiologic and clinical entity because of its etiologic and therapeutic peculiarities. Although it frequently occurs in the setting of a prolonged QT interval due to hypokalemia or antiarrhythmic drugs, it can also occur without these transient abnormalities. In the present study we investigated the electrophysiologic characteristics of this arrhythmia in patients without transient precipitating factors.

Patients

The study population included 21 patients, 13 males and eight females, ages 30–72 years, in whom torsades de pointes was induced during electrophysiologic studies or occurred spontaneously (table 1). All but two patients had had at least three spontaneous episodes of torsades de pointes documented electrocardiographically, either by computer-monitored continuous electrocardiographic recording in a coronary care unit or Holter monitoring during observation periods of 4–13 days. The two patients in whom torsades de pointes occurred only during electrophysiologic studies had frequent multiform premature ventricular depolarizations and nonsustained ventricular tachycardia, and thus did not fulfill strict criteria for the diagnosis of torsades de pointes. Eleven patients had suffered cardiac arrest, with ventricular tachycardia or fibrillation documented as the mechanism. Torsades de pointes always occurred in the absence of neonatal asphyxia and intraventricular hemorrhage by Doppler ultrasound. J Pediatr 95: 775, 1979


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of bradycardias, antiarrhythmic or digitalis therapy, electrolyte imbalance, and recent (less than 4 weeks) myocardial infarction. For this analysis we have excluded patients in whom torsades de pointes was observed only during transient abnormalities of rhythm, metabolic state or therapy with agents that have been reported to induce this arrhythmia.

The cardiac diagnoses and other demographic data are presented in Table 1. The QT/QTc intervals were measured during sinus rhythm when the patients were having spontaneous episodes. With the Bazett method, 14 patients had normal QTc intervals and seven had minimally prolonged QTc intervals (greater than 0.392 for males and greater than 0.44 for fe-

**Table 1. Demographic Data**

<table>
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<tr>
<th>Pt.</th>
<th>Age (years)</th>
<th>Sex</th>
<th>Diagnosis</th>
<th>ECG morphology</th>
<th>QT/QTc (sec)</th>
<th>Clinical arrhythmia</th>
<th>Number of complexes in paroxysms of torsades de pointes</th>
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<td>F</td>
<td>CAD, Vent. an.</td>
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<td>TdP, VF</td>
<td>6-26</td>
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<td>TdP</td>
<td>6-12</td>
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<td>VPDs</td>
<td>15-23</td>
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<td>AS, CAD</td>
<td>AMI, LVI</td>
<td>0.31/0.37</td>
<td>TdP, VF</td>
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<td>AMI</td>
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<td>TdP, VF</td>
<td>9-24</td>
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<tr>
<td>6</td>
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<td>M</td>
<td>CAD</td>
<td>WNL</td>
<td>0.38/0.38</td>
<td>TdP, VF</td>
<td>12-19</td>
</tr>
<tr>
<td>7</td>
<td>59</td>
<td>F</td>
<td>CM</td>
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<td>TdP</td>
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<td>62</td>
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<td>55</td>
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<td>0.39/0.30</td>
<td>TdP</td>
<td>6-34</td>
</tr>
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<td>TdP, VF</td>
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</table>

**Abbreviations:** AF = atrial fibrillation; AMI = anterior myocardial infarction; AS = aortic stenosis; CAD = coronary artery disease; CM = cardiomyopathy; HTN = hypertension; IMI = inferior myocardial infarction; LAHB = left anterior hemiblock; LBBB = left bundle branch block; LVH = left ventricular hypertrophy; MVP = mitral valve prolapse; NSSTTWA = nonspecific ST- and T-wave abnormalities; PEHD = primary electrical heart disease; QTc = measured in control state by Bazett's method; RHD = rheumatic heart disease; RV = right ventricle; TdP = torsades de pointes; Vent. an. = ventricular aneurysm; VF = ventricular fibrillation; VPD = ventricular premature depolarization; WNL = within normal limits.
two patients had prolonged QT intervals corrected for rate. No patient had a QTc interval greater than 0.51 second. Atrial pacing did not cause an increase in QT or QTc in any patient.

Methods

All patients were referred specifically for evaluation of ventricular tachycardia or syncope. The diagnosis of torsades de pointes was made using the electrophysiographic criteria suggested by Dessertenne,1 Slama et al.,7,11 Krikler and Curry12 and the WHO/ISFS Task Force.14 These criteria include (1) paroxysms of ventricular tachycardia, during which the rhythm is irregular and usually has an average rate of 200–250 beats/min; (2) progressive changes in amplitude and polarity of the QRS complexes, usually in an interval of five to 10 complexes such that the QRS axis changes and the complexes appear to be twisting around the isoelectric baseline; (3) spontaneous termination after a few complexes or as long as several minutes and consecutive episodes that vary widely in morphology and duration (fig. 1), (4) occasional progression of the arrhythmia to sustained uniform morphology ventricular tachycardia or ventricular fibrillation. We chose six complexes as the minimum number of complexes to define torsades de pointes.

Electrophysiologic studies were performed in the postabsorptive state after written informed consent had been obtained. All antiarrhythmic agents had been discontinued for more than five half-lives before study and plasma concentrations at the time of the study were below the measurable level by routine laboratory methods. Clinical status was stable and electrolyte measurements were normal and stable. Two to four electrode catheters were inserted percutaneously or by cutdown and positioned in the heart under fluoroscopic guidance. In each patient, catheters were positioned in at least one atrial site, usually the high right atrium, the His bundle position and the right ventricular apex. In selected patients, catheters were positioned in the right ventricular outflow tract (six patients) or left ventricle (seven patients) or both (five patients). Left ventricular electrograms were recorded in 18 of 21 patients, either from a left ventricular or coronary sinus catheter. Standard quadrapololar electrode catheters were used when recording and pacing from the same site were required. Stimulation was performed using a specially designed programmable stimulator and optically isolated constant current sources (Bloom Associates, Ltd.). The stimuli were rectangular pulses 1 msec in duration delivered at twice diastolic threshold (0.5–2.5 mA). Leakage current was regularly monitored and never exceeded 6 μA.

The protocol of programmed stimulation included: atrial pacing at incremental rates (100–250 beats/min), premature atrial stimulation during sinus rhythm and/or atrial pacing, ventricular pacing at incremental rates (100–250 beats/min), or premature ventricular stimulation during sinus rhythm and/or ventricular pacing. Single, double and triple extrastimuli were used. Diastole was scanned with the first premature stimulus (S1) in 10–20-msec decrements until ventricular refractoriness. If no ventricular arrhythmia was initiated, the first premature stimulus (S2) was introduced at a coupling interval (S1S2) 50–100 msec longer than the ventricular effective refractory period. A second premature stimulus (S3) was introduced at an interval (S2S3) equal to S1S2. S3S3 was shortened by 10–20 msec until S3 failed to capture the ventricle or initiate the arrhythmia. The S3S3 was then decreased until S3 again captured the ventricle. The S3S3 was then shortened again. This procedure was continued until both S2 and S3 reached refractoriness. If no ventricular arrhythmia was initiated, a third premature stimulus was introduced at an interval (S1S3) equal to S1S2. The S3S3 was again programmed to 50–100 msec greater than the ventricular effective refractory period and S3S3 and S3S3, which were initially equal to S1S2, were decreased sequentially until S3 and S3 did not produce an evoked response. Stimulation was performed at two or more cycle lengths in 19 patients and at the right ventricular apex in all patients. Stimulation was also performed at the right ventricular outflow tract in six patients and in the left ventricle in seven patients if the arrhythmia was not initiated by pacing at the right ventricular apex. If sustained ventricular tachycardia was initiated, single, double and triple extrastimuli or rapid ventricular pacing (200–300 beats/min) was used in an attempt to terminate the arrhythmia.

Immediately after baseline studies, in which torsades de pointes was initiated at least three times, procainamide (1–2 g; 50 mg/min) or quinidine (600–1200 mg; 20 mg/min) was infused intravenously and programmed stimulation was repeated after the infusion.

Intracardiac recordings were filtered at 30–500 Hz and displayed simultaneously with two or three electrocardiographic leads and 10-msec time lines on an oscillographic recorder (Electronics for Medicine VR16). All data were stored on magnetic analog tape and later retrieved on photographic paper at speeds of 75–200 mm/sec.

Results

Initiation of Torsades de Pointes

Torsades de pointes was initiated by programmed ventricular stimulation in 19 of the 21 patients during control studies in the absence of antiarrhythmic drugs. The arrhythmia was initiated during both sinus rhythm and ventricular pacing in two patients and during ventricular pacing only in 17 patients (fig. 2). Initiation was accomplished by a single extrastimulus in two patients and required double extrastimuli in 17. Left ventricular stimulation was necessary in one patient. Rapid ventricular pacing also induced torsades de pointes in two of the 19 patients. In all 19 patients, initiation was reproducible; it was ac-
FIGURE 2. Initiation of torsades de pointes during normal sinus rhythm and ventricular pacing in patient 9. ECG leads 1, 2 and V1 are shown with electrograms recorded in the coronary sinus (CS), His bundle area (HBE) and right ventricular apex (RVA) with 100-msec time marks (T). (top panel) Two premature stimuli (arrows) were introduced after the second sinus complex and a paroxysm of torsades de pointes was initiated. (bottom panel) Two premature stimuli (arrows) delivered after the third paced complex initiated a typical episode of torsades de pointes, which terminated spontaneously.

In patients 7 and 21, in whom torsades de pointes occurred spontaneously, the arrhythmia could not be initiated by programmed stimulation. In these patients, single, double and triple extrastimuli introduced during sinus rhythm and ventricular pacing at two cycle lengths and rapid ventricular pacing were performed at the right ventricular apex, right ventricular outflow tract and at least two left ventricular sites. No clinical or electrophysiologic measurements distinguished these two patients from the 19 patients in whom torsades de pointes was inducible.

Observations During Torsades de Pointes

The QRS morphology during the induced paroxysms of torsades de pointes showed the typical undulating pattern of the spontaneous episodes. The duration, rate and electrocardiographic patterns of the induced episodes were not different from those of the spontaneous ones (compare figures 1 and 2). The self-terminating paroxysms of induced torsades de pointes lasted a minimum of six complexes and a maximum of 34 complexes. The mean duration of these paroxysms was 17 ± 12 complexes. The duration and morphology of episodes varied widely in each patient during consecutive initiations. Paroxysms of torsades de pointes initiated during normal sinus rhythm tended to be shorter and have a longer mean cycle length than those initiated during ventricular pacing (fig. 2). Similarly, consecutive spontaneous episodes varied widely in duration and morphology.

The RR intervals and interelectrogram intervals measured at the site of stimulation varied widely during sequential paroxysms in individual patients. The shortest interelectrogram interval in the group was 110 msec and the longest was 360 msec. During individual paroxysms, the difference between the shortest and longest interelectrogram interval varied from 50–220 msec. The mean interelectrogram interval for all paroxysms in this group was 258 ± 87 msec. Although the interelectrogram intervals tended to lengthen and shorten sequentially in an oscillatory pattern, no specific pattern of oscillation was iden-
Termination of Torsades de Pointes

One hundred forty-eight of 155 episodes of torsades de pointes initiated in these patients terminated spontaneously. In 97 of 148 episodes, termination occurred after a sequence of three to five complexes showing progressively increasing interelectrogram intervals (fig. 2, bottom panel). In 22 episodes, spontaneous termination followed acceleration of the local electrograms and in 29, no specific pattern of interelectrogram intervals appeared. All episodes of torsades de pointes were too short to attempt termination by programmed stimulation.

Spontaneous Conversion of Torsades de Pointes to Sustained Ventricular Tachyarrhythmia

In four of 155 episodes (three patients), torsades de pointes progressed to sustained ventricular fibrillation, which was terminated by countershock. In these patients (patients 5, 15 and 20), cardiac arrest due to electrocardiographically documented ventricular fibrillation had occurred before study. Except for degeneration to ventricular fibrillation, the characteristics (e.g., rate, RR interval variability) of these episodes were not significantly different from those that terminated spontaneously.

In three of 155 episodes (three patients), torsades de pointes organized into a sustained ventricular tachycardia of uniform morphology or ventricular flutter (fig. 3). The cycle lengths of these tachycardias were 170, 190 and 205 msec. When the uniform morphology developed, the activation pattern observed in the ventricular electrograms became constant. Each of these ventricular tachycardias required DC cardioversion because they rapidly produced hemodynamic deterioration.

Response to Procainamide or Quinidine

In eight patients the stimulation protocol that produced torsades de pointes during control studies initiated a sustained ventricular tachycardia with a uniform morphology after administration of procainamide (five patients) or quinidine (three patients) (fig. 4). The cycle length of these tachycardias ranged from 250–420 msec and had electrophysiologic
characteristics identical to those described for recurrent sustained ventricular tachycardia. In six patients, this tachycardia could be terminated by programmed extrastimuli (four patients) or rapid ventricular pacing (two patients) (Fig. 4). Termination required cardioversion in the two remaining patients. Torsades de pointes was the only inducible ventricular arrhythmia during the control studies. In none was sustained ventricular tachycardia with uniform morphology inducible before procainamide or quinidine.

In two of the eight patients, ventricular tachycardia or torsades de pointes was inducible or occurred spontaneously during treatment with all standard antiarrhythmic drugs. In these two patients, surgery was performed for ablation of arrhythmias using intraoperative mapping to localize the site of origin. The surgery was performed during treatment with procainamide to maintain a uniform-morphology ventricular tachycardia. In each, the site of origin was identified and resected. In neither patient was torsades de pointes or sustained uniform-morphology ventricular tachycardia inducible by programmed electrical stimulation postoperatively, nor have spontaneous episodes occurred in either patient.

In nine patients the acute administration of procainamide or quinidine prevented initiation of torsades de pointes and any other ventricular tachyarrhythmia. In four other patients the initiation and characteristics of torsades de pointes were unaffected by the administration of procainamide (four patients) or quinidine (three patients).

Discussion

The occurrence of torsades de pointes in clinically stable patients without electrolyte disturbance or antiarrhythmic medication and normal or minimally prolonged QTc intervals has been reported but has been infrequently emphasized.

Relationship Between Spontaneous and Induced Torsades de Pointes

The characteristics of the induced torsades de pointes were similar to those that occurred spontaneously. Although the QRS morphology and duration of individual paroxysms of torsades de pointes varied widely between consecutive episodes, in each patient the induced episodes behaved similarly to spontaneous episodes. In three patients, torsades de pointes spontaneously converted to sustained, uniform-morphology ventricular tachycardia; in each of these patients, spontaneous episodes of rapid sustained, uniform-morphology ventricular tachycardia or ventricular flutter had occurred. Furthermore, the patients in whom induced torsades de pointes progressed to ventricular fibrillation similarly had recurrent clinical episodes of ventricular fibrillation that produced cardiac arrest.
Electrophysiologic Characteristics of Torsades de Pointes

The torsades de pointes investigated in this series met the most frequently used criteria for a reentrant arrhythmia, namely, reproducible initiation by premature stimulation. In 19 of 21 patients (92%), the torsades de pointes was inducible by one or two ventricular extrastimuli. These data are in agreement with the studies of Evans et al., in which torsades de pointes was inducible by programmed electrical stimulation in four of four patients.

Most paroxysms of torsades de pointes terminated spontaneously. This observation does not provide data regarding the mechanism of the arrhythmia because it is compatible with either an automatic or reentrant arrhythmia. The spontaneous evolution of torsades de pointes into a uniform-morphology ventricular tachycardia, which has the typical characteristics of a reentrant ventricular tachycardia being terminated by programmed stimulation and/or cardioversion, however, does support the reentrant mechanism.

Effect of Procainamide and Quinidine

The conversion of torsades de pointes to typical sustained ventricular tachycardia by administration of either procainamide or quinidine supports the reentrant-mechanism hypothesis. The resultant ventricular tachycardias have the characteristic response to programmed stimulation generally reported in reentrant ventricular tachycardia. Similar clinical responses to procainamide have been reported. Furthermore, cure of torsades de pointes was accomplished by resection of the site of origin of this typical ventricular tachycardia in two patients, suggesting that the torsades de pointes and the uniform-morphology ventricular tachycardia were actually different manifestations of the same arrhythmia.

Unifying Hypothesis Regarding the Mechanism of Recurrent Ventricular Tachyarrhythmias

Recurrent sustained ventricular tachycardia, a regular and sustained tachyarrhythmia, usually fulfills the clinical electrophysiologic characteristics that suggest a reentrant mechanism. Recurrent ventricular fibrillation occurring in clinically stable patients can also, in certain patients, be initiated by programmed stimulation, suggesting initiation by reentry. Although ventricular electrograms during ventricular fibrillation tend to be irregular and fractionated, local ventricular electrical activity at the onset and during ventricular fibrillation can be regular and discrete. The relationship of these recurrent ventricular tachyarrhythmias is unclear; however, a better understanding of this relationship may be obtained by study of torsades de pointes, a variant type of ventricular tachycardia that has characteristics of both of these classic ventricular tachyarrhythmias.

Evans et al. and Fontaine et al. presented data that support a reentrant mechanism for this arrhythmia. Evans et al. showed that torsades de pointes was inducible in four patients. Fontaine et al. suggested that torsades de pointes results from a reentrant mechanism because their patients had prolonged ventricular refractory periods and delayed potentials recorded during normal sinus rhythm.

Our data suggest that torsades de pointes may be a variant form of sustained ventricular tachycardia. Torsades de pointes may be a rapid and, at its origin, a regular ventricular tachyarrhythmia in which the electrocardiographic irregularity is caused by variable patterns of ventricular activation caused by changing exit sites from the origin of the arrhythmia or local conduction block related to encountering refractoriness as a result of the short cycle length of the tachycardia. Moreover, torsades de pointes can spontaneously regularize into a uniform-morphology tachyarrhythmia as refractoriness decreases with the short cycle length, eliminating local block, or as ventricular activation proceeds via a single exit site as functional changes at the other exit sites preclude exit from them. Further support of this hypothesis is the observation that procainamide or quinidine, which increase the cycle length of ventricular tachycardia, frequently convert this irregular tachyarrhythmia to a ventricular tachyarrhythmia with a uniform morphology that has characteristics typical of reentrant ventricular tachycardia. The mechanism for this conversion may be a relatively greater depression of conduction than increase in refractoriness caused by procainamide or quinidine.

These observations and hypotheses are consistent with studies in experimental chronic infarction in which arrhythmias morphologically similar to torsades de pointes are caused by rapid regular ventricular tachyarrhythmias that produce variable ventricular activation patterns. Nhon et al. showed that varying exit sites from a rapid ventricular tachycardia can produce a pattern of torsades de pointes. They suggested that varying exit sites or the course of the reentrant circuit could produce the QRS morphology that is typical of this arrhythmia. Ideker et al. showed that the electrocardiographic pattern of torsades de pointes can also be caused by the successive summation of two to four depolarization wave fronts as they course over the epicardium from the site of origin of a rapid ventricular tachycardia. The random and continuously varying summation of these wave fronts produces undulating cyclical QRS patterns.

Torsades de pointes, on the other hand, can also lead to ventricular fibrillation, presumably by the same mechanism by which rapid uniform-morphology ventricular tachycardia or rapid ventricular pacing produces ventricular fibrillation. Ventricular activation can become progressively chaotic as wave fronts fractionate and encounter areas of differing refractoriness and conductivity. This heterogeneous pattern of conduction and recovery can lead to the self-sustaining fibrillatory process.

Other explanations are possible. Multifocal ventricular tachycardia with fusion complexes producing the varying QRS morphologies might also explain our observations; however, we believe our hypothesis is the best explanation.
Relationship to Other Investigations

The electrocardiographic features of torsades de pointes were reported over half a century ago by MacWilliams and Wiggers. Schwartz et al. thoroughly described the syndrome, including the association with bradycardia and prolonged QT intervals. They also emphasized the interrelationship between this arrhythmia and uniform-morphology ventricular tachycardia.3 They termed ventricular fibrillation,3 recurrent ventricular fibrillation,3 and pseudofibrillation,3 cardiac flutter,4 atypical ventricular tachycardia,5 ventricular fibrillo-flutter,6 polymorphous ventricular tachycardia,7 and torsades de pointes.1 This surfeit of names suggests a paucity of data regarding the cause of this arrhythmia.

We excluded from analysis patients in whom torsades de pointes was related to transient factors, such as hypokalemia or antiarrhythmic drug therapy. Twenty of our 21 patients had chronic and stable structural heart disease, and 12 (60%) had chronic coronary artery disease. Most investigators have reported a similar incidence of organic cardiac disease. Although torsades de pointes has been reported to occur in patients with stable chronic heart disease, and particularly coronary artery disease, this etiology has not been emphasized. We cannot comment on the relative frequency of the different etiologies of torsades de pointes because we reported only patients in whom transient factors did not play a role. The patients in whom torsades de pointes is caused by transient pharmacologic or metabolic abnormalities are less stable clinically and more difficult to study. Furthermore, these patients are usually promptly treated by removing the underlying cause and diagnostic electrophysiologic catheterization is necessary less often. Our present studies, therefore, have been confined to patients in whom the arrhythmia occurred in a stable setting.

The arrhythmias in both groups, however, are morphologically similar and, in the final analysis, arrhythmias are named for their morphology rather than etiology. Although some may disagree with calling the arrhythmia we describe torsades de pointes, because QT interval prolongation was not always present, the arrhythmias in our patients met the morphologic criteria suggested by most authors. Until further data are available, we believe greater understanding of these ventricular tachyarrhythmias will be gained by considering them in the same group of arrhythmias rather than creating many categories. Clarification of the nosographic confusion surrounding this arrhythmia is beyond the scope of the present work; however, because these arrhythmias are so similar in morphology, etiology and clinical significance, we suggest that greater uniformity in terminology may allow an organized approach to investigating their mechanisms.

Our data and those of others suggest that torsades de pointes in the setting of organic heart disease is a reentrant tachyarrhythmia with electrophysiologic characteristics of both typical ventricular tachycardia and ventricular fibrillation. Further study of the anatomic and electrophysiologic substrate of these three groups of ventricular tachyarrhythmias may produce greater understanding of the causes and processes that result in sudden cardiac death.

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