First introduced as a new clinical entity in 1992, the Brugada syndrome is associated with a relatively high risk of sudden death in young adults, and occasionally in children and infants. Recent years have witnessed a striking proliferation of papers dealing with the clinical and basic aspects of the disease. Characterized by a coved-type ST-segment elevation in the right precordial leads of the electrocardiogram (ECG), the Brugada syndrome has a genetic basis that thus far has been linked only to mutations in SCN5A, the gene that encodes the α-subunit of the sodium channel. The Brugada ECG is often concealed, but can be unmasked or modulated by a number of drugs and pathophysiological states including sodium channel blockers, a febrile state, vagotonic agents, tricyclic antidepressants, as well as cocaine and propranolol intoxication. Average age at the time of initial diagnosis or sudden death is 40 ± 22, with the youngest patient diagnosed at 2 days of age and the oldest at 84 years. This review provides an overview of the clinical, genetic, molecular, and cellular aspects of the Brugada syndrome, incorporating the results of two recent consensus conferences. Controversies with regard to risk stratification and newly proposed pharmacologic strategies are discussed. (PACE 2006; 29:1130–1159)

ventricular arrhythmias, sudden cardiac death, atrial fibrillation, ventricular tachycardia, ventricular fibrillation, inherited disease

Introduction

Since its introduction as a new clinical entity by Pedro and Josep Brugada in 1992,1 the Brugada syndrome has attracted great interest because of its high incidence in many parts of the world and its association with high risk of sudden death, especially in males as they enter their third and fourth decades of life. Recent years have witnessed a dramatic rise in the number of reported cases and a great proliferation of papers serving to define the clinical, genetic, cellular, ionic, and molecular aspects of the disease.2 A consensus conference report published in 2002 delineated diagnostic criteria for the syndrome.3,4 A second consensus conference report published in 2005 focused on risk stratification schemes and approaches to therapy.5,6 This review provides an in-depth overview of the clinical, genetic, molecular, and cellular aspects of the Brugada syndrome, incorporating the results of the two consensus conference reports, and the numerous clinical and basic publications on the subject. Controversies with regard to the involvement of anatomic structural defects, the approach to risk stratification, and new data relative to proposed pharmacologic strategies are discussed in detail. The proposed terminology, diagnostic criteria, risk stratification schemes and device, and pharmacologic approach to therapy discussed are based on available clinical and basic studies, and should be considered a work-in-progress that will require significant fine-tuning as confirmatory data from molecular studies and prospective trials become available.

Clinical Characteristics and Diagnostic Criteria

Characterized by an ST-segment elevation in the right precordial electrocardiogram (ECG) leads and a high incidence of sudden death in patients with structurally normal hearts, the Brugada syndrome generally manifests during adulthood. The average age at the time of initial diagnosis or sudden death is 40 ± 22. The youngest patient diagnosed with the syndrome is 2 days of age, and the oldest is 84 years.

A recent report by Skinner et al. provides a direct link between sudden infant death syndrome and the Brugada syndrome.7 This 19-day-old infant was successfully resuscitated from ventricular fibrillation (VF). Although the basal ECG did not show features of Brugada syndrome, DNA analysis revealed a missense mutation (R1193Q) in SCN5A, previously linked with the Brugada syndrome.

Because the ECG is so dynamic and often concealed, it is difficult to estimate the true prevalence of the disease in the general population.8 The prevalence of the Brugada syndrome is estimated at 1–5 per 10,000 inhabitants worldwide. The frequency is lower in western countries and higher (≥5 per 10,000) in Southeast Asia, especially in Thailand and the Philippines where Brugada syndrome is considered to be the major cause of sudden death in young adults.
of sudden death in young individuals. In these
countries, the syndrome is often referred to as
sudden unexplained nocturnal death syndrome
(SUNDS).9,10

The electrocardiographic manifestations of
the Brugada syndrome when concealed can be
unmasked by sodium channel blockers, a febrile
state, or vagotonic agents.11–14 Three types of re-
polarization patterns in the right precordial leads
are recognized (Table I and Fig. 1).3,4 Type 1 ST-
segment elevation is diagnostic of Brugada syn-
drome and is characterized by a coved ST-segment
elevation \( \geq 2 \text{ mm} \) (0.2 mV) followed by a negative
T wave. Type 2 ST-segment elevation has a saddle-
back appearance with a high take-off ST-segment
elevation \( \geq 2 \text{ mm} \) followed by a trough display-
ing \( \geq 1 \text{ mm} \) ST elevation followed by either a posi-
tive or biphasic T wave. Type 3 ST-segment eleva-
tion has either a saddleback or coved appearance
with an ST-segment elevation of \( < 1 \text{ mm} \). These
three patterns may be observed sequentially in the
same patient or following the introduction of spe-
cific drugs. Type 2 and 3 ST-segment elevation
should not be considered diagnostic of the Bru-
gada syndrome. A Brugada ECG refers to the man-
ifestation of a Type 1 ST-segment elevation. Bru-
gada syndrome is definitively diagnosed when a
Type 1 ST-segment elevation (Brugada ECG) is ob-
served in more than one right precordial lead (V1-
V3), in the presence or absence of sodium chan-
nel blocking agent, and in conjunction with one or
more of the following: documented VF, polymor-
phic ventricular tachycardia (VT); a family history
of SCD (<45 years old); coved-type ECGs in fam-
ily members; inducibility of VT with programmed
electrical stimulation (PES); syncpe; or nocturnal
agonal respiration.3–6

Differential diagnosis of the Brugada syn-
drome must be approached with care since ST-
segment elevation is associated with a wide variety
of benign as well as malignant pathophysio-
logic conditions. Definitive diagnosis is difficult
when the degree of basal ST-segment elevation
is relatively small and the specificity of sodium
channel blockers, such as flecainide, ajmaline,
procainamide, disopyramide, propafenone, and

<table>
<thead>
<tr>
<th>J-point</th>
<th>Type 1</th>
<th>Type 2</th>
<th>Type 3</th>
</tr>
</thead>
<tbody>
<tr>
<td>≥2 mm</td>
<td>≥2 mm</td>
<td>≥2 mm</td>
<td></td>
</tr>
<tr>
<td>T-wave</td>
<td>Negative</td>
<td>Positive or biphasic</td>
<td>Positive</td>
</tr>
<tr>
<td>ST-T configuration</td>
<td>Coved type</td>
<td>Saddleback</td>
<td>Saddleback</td>
</tr>
<tr>
<td>ST segment (terminal portion)</td>
<td>Gradually descending</td>
<td>Elevated ( \geq 1 \text{ mm} )</td>
<td>Elevated ( &lt; 1 \text{ mm} )</td>
</tr>
</tbody>
</table>

\( 1 \text{ mm} = 0.1 \text{ mV} \), the terminal portion of the ST segment refers to the latter half of the ST segment. From Wilde et al.3 with permission.

Figure 1. Three Types of ST-segment elevation generally ob-
erved in patients with the Brugada syndrome. Shown are precordial
leads recorded from a patient diag-
nosed with the Brugada syndrome.
Note the dynamic ECG changes oc-
curring over a period of 2 days.
The left panel shows a clear Type
1 ECG, which is diagnostic of the
Brugada syndrome. A saddleback
ST-segment elevation (Type 2) is
observed on February 7, 1999. The
ST segment is further normalized
on February 13, 1999, showing a
Type 3 ECG. Modified from Wilde
et al.4 with permission.
Table II.

Drugs Used to Unmask the Brugada Syndrome

<table>
<thead>
<tr>
<th>Drug</th>
<th>Dosage</th>
<th>Route</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ajmaline</td>
<td>1 mg/kg/5 min, i.v.</td>
<td></td>
</tr>
<tr>
<td>Flecainide</td>
<td>2 mg/kg/10 min, i.v. (400 mg, p.o.)</td>
<td></td>
</tr>
<tr>
<td>Procainamide</td>
<td>10 mg/kg/10 min, i.v.</td>
<td></td>
</tr>
<tr>
<td>Pilsicainide</td>
<td>1 mg/kg/10 min, i.v.</td>
<td></td>
</tr>
</tbody>
</table>

pilsicainide\(^{12,15,16}\) to identify patients at risk is uncertain. The recommended dosage regimens for the most frequently used agents are listed in Table II. A comparison of intravenous ajmaline and flecainide in the same cohort of patients revealed that ajmaline is more effective in unmasking the syndrome.\(^{17}\) Flecainide failed in 7 of 22 cases (32%) unmasked by ajmaline. A greater inhibition of I\(_v\) by flecainide renders it less effective than ajmaline.

The sodium challenge should be monitored with a continuous ECG recording (speed 10 mm/s and interposed 50 mm/s) and should be terminated when (1) the diagnostic Type 1 ST-segment elevation, or Brugada ECG, develops; (2) ST segment in Type 2 increases by ≥2 mm; (3) premature ventricular beats or other arrhythmias develop; or (4) QRS widens to ≥130% of baseline. Sodium channel blockers should be used with particular caution in the presence of atrial and/or ventricular conduction disease (presence of wide QRS, wide P waves, or prolonged PR intervals). Isolated cases of mechano-electrical dissociation have been reported. Isoproterenol and sodium lactate may be effective antidotes.\(^{5,6}\)

Confounding factor(s) that could account for the ECG abnormality need to be carefully excluded (Table III). Exaggerated ST-segment elevation is sometimes observed for a brief period following DC cardioversion, and this must be factored in when first evaluating a patient following defibrillation.\(^{18}\)

Another important confounding factor is the ST-elevation encountered in well-trained athletes. The ST-segment elevation encountered in athletes can be distinguished by virtue of the fact that it is upsloping rather than downsloping and is largely unaffected by challenge with a sodium channel blocker. In addition, a variety of drugs have been reported to produce a Brugada-like ST-segment elevation (Table IV). Myocarditis and some forms of arrhythmogenic right ventricular cardiomyopathy/dysplasia (ARVC/D) can lead to a Brugada-like phenotype (see below). The extent to which these acquired forms of Brugada syndromes have a genetic predisposition is the subject of intense investigation.

Diagnosis of Brugada syndrome is also considered positive when a Type 2 (saddleback pattern) or Type 3 ST-segment elevation is observed in more than one right precordial lead under baseline conditions and can be converted to the diagnostic Type 1 pattern occurs upon exposure to sodium channel blocker (ST-segment elevation should be ≥2 mm). One or more of the clinical criteria described above should also be present. Drug-induced conversion of Type 3 to Type 2 ST-segment elevation is considered inconclusive for diagnosis of Brugada syndrome.

Most cases of Brugada syndrome display right precordial ST-segment elevation, although isolated cases of inferior lead\(^{19,20}\) or left precordial lead\(^{21}\) ST-segment elevation have been reported in Brugada-like syndromes, in some cases associated with SCN5A mutations. In rare cases, ST-segment elevation is observed in all precordial leads (unpublished observation).

Placement of the right precordial leads in a superior position (up to the 2\(^{nd}\) intercostal spaces above normal) can increase the sensitivity of the ECG for detecting the Brugada phenotype in some patients, both in the presence or absence of a drug challenge (Fig. 2).\(^{22,23}\) Studies are underway to
Table IV. Drug-Induced Brugada-Like ECG Patterns

<table>
<thead>
<tr>
<th>Category</th>
<th>Drugs</th>
</tr>
</thead>
</table>
| Antiarrhythmic drugs | Na\(^+\) channel blockers  
Class IC drugs (Flecainide\(^ {12,15,142,207,208}\)  
Pilsicainide\(^ {146,209}\) Propafenone\(^ {210}\))  
Class IA drugs (Ajmaline\(^ {12,211}\) Procainamide\(^ {12,13}\)  
Disopyramide\(^ {4,13}\) Cibenzoline\(^ {212,213}\))  
Ca\(^{2+}\) channel blockers  
Verapamil  
\(\beta\)-Blockers  
Propranolol intoxication\(^ {214}\) |
| Antianginal drugs | Ca\(^{2+}\) channel blockers  
Nifedipine, diltiazem  
Nitrate  
Isosorbide dinitrate, nitroglycerine\(^ {215}\)  
K\(^+\) channel openers  
Nicorandil |
| Psychotropic drugs | Tricyclic antidepressants\(^ {216}\)  
Amitriptyline\(^ {217,218}\) Nortriptyline,\(^ {151}\) Desipramine\(^ {149}\)  
Clomipramine\(^ {150}\)  
Tetracyclic antidepressants  
Maprotiline\(^ {217}\)  
Phenothiazine  
Perphenazine\(^ {217}\) Cyamemazine.  
Selective serotonin reuptake inhibitors  
Fluoxetine\(^ {218}\)  
Lithium\(^ {157}\) |
| Other drugs | Histaminic H1 receptor antagonists  
Dimenhydrinate\(^ {152}\)  
Diphenhydramine\(^ {219}\)  
Cocaine intoxication\(^ {153,220}\)  
Alcohol intoxication |

Modified from Antzelevitch et al.\(^ {5,6}\) and Shimizu\(^ {221}\) with permission.

ascertain whether the greater sensitivity is at the cost of a lower specificity and whether a Type I ECG in the elevated leads is as predictive of events as a Type I ECG in standard leads. A recent report demonstrates that as many as 1.3% of normal Korean males display a Type 2, but not Type 1, ST-segment elevation when the right precordial leads are recorded from a superior position.\(^ {24}\)

A slight prolongation of the QT interval is sometimes observed associated with the ST-segment elevation.\(^ {16,25,26}\) The QT interval is prolonged more in the right versus left precordial leads, presumably due to a preferential prolongation of action potential duration (APD) in right ventricular (RV) epicardium secondary to accentuation of the action potential (AP) notch.\(^ {27}\) A corrected QT (QTc) >460 ms in V2 has been shown to be associated with arrhythmic risk.\(^ {28}\) Depolarization abnormalities including prolongation of

Figure 2. Shift of right precordial leads to 2\(^{nd}\) and 3\(^{rd}\) intercostal space unmasks a Type 1 Brugada ECG. (Top) Plot of 87 unipolar electrode sites (dots) and of 6 precordial electrocardiograms (ECG) (crosses). Eighty-seven lead points are arranged in a lattice-like pattern (13 × 7 matrix), except for four lead points on both mid-axillary lines and covered the entire thoracic surface. V1 and V2 leads of the ECG are located between D5 and E5, and between E5 and F5, respectively, whereas V4, V5, and V6 are coincident with G4, H4, and I4, respectively. (Bottom) Twelve-leads electrocardiograms (ECG) in a patient with Brugada syndrome. Type 2 saddleback ST-segment elevation was observed in V1 and V2 of the standard 12-lead ECG (4\(^{th}\) intercostal space), whereas typical Type 1 coved-type ST-segment elevation was apparent in V1 and V2 recorded from the 2\(^{nd}\) and 3\(^{rd}\) intercostal space (arrows). Modified from Antzelevitch et al.\(^ {5,6}\) with permission.
P-wave duration, PR and QRS intervals are frequently observed, particularly in patients linked to SCN5A mutations. PR prolongation likely reflects HV conduction delay.

More recent reports indicate that approximately 20% of Brugada syndrome patients develop supraventricular arrhythmias. It is as yet unknown whether atrial vulnerability is correlated with ventricular inducibility of arrhythmias or whether atrial arrhythmias may serve as triggering events for VT/VF, although the latter seems unlikely based on current knowledge. Slowed atrial conduction as well as atrial standstill have been reported in association with the syndrome.

In many cases, arrhythmia initiation is bradycardia-related. This may contribute to the higher incidence of sudden death at night in individuals with the syndrome and may account for the success of pacing in controlling the arrhythmia in isolated cases of the syndrome. However, not all patients die at night and not all the cases are controlled with rapid ventricular pacing. South Asian patients who have the ECG pattern usually develop VT/VF during sleep at night. Makiyama and co-workers reported that loss-of-function SCN5A mutations resulting in Brugada syndrome are distinguished by profound bradycardairrhythmias. Pertinent to this observation is the recent report by Scornik and co-workers demonstrating expression of the cardiac sodium channel gene, SCN5A, in intracardiac ganglia. This interesting finding suggests that loss-of-function mutations in SCN5A may not only create the substrate for reentry in ventricular myocardium, but may also increase vagal activity in intracardiac ganglia, thus facilitating the development of arrhythmias in patients with the Brugada syndrome.

A polymorphic VT resembling a rapid Torsade de Pointes (TdP) arrhythmia is most commonly associated with the Brugada syndrome. Monomorphic VT is observed infrequently and is generally more prevalent in children and infants. VT/VF often terminates spontaneously in patients with the Brugada syndrome, as first reported by Bjørregaard et al. This may explain why patients wake up at night after episodes of agonal respiration caused by the arrhythmia.

SUNDS, also known as SUDS, a disorder most prevalent in Southeast Asia, and Brugada syndrome have recently been shown to be phenotypically, genetically, and functionally the same disorder. Sudden and unexpected death of young adults during sleep, known in the Philippines as bangungut (“to rise and moan in sleep”), was first described in the Philippine medical literature in 1917. In Japan, this syndrome, known as pokkuri (“sudden and unexpectedly ceased phenomena”), was reported as early as 1959. In 1997, Nade-Manee et al. reported that among 27 Thai men referred for aborted cases of what was known in Thailand as Lai Tai (“death during sleep”), as many as 16 had the ECG pattern of Brugada Syndrome. In their review of the literature in 1999, Alings and Wilde found that of the 163 patients who met the criteria for Brugada Syndrome, 58% were of Asian origin.

**Relationship with Structural Heart Disease**

This issue has been the subject of much debate and one that is in a rapid course of evolution. Like the long-QT and short-QT syndromes, Brugada syndrome is traditionally thought of as a primary electrical cardiac disease arising in myocardium that is otherwise structurally normal. Although cardiac function may seem normal on noninvasive examination using echocardiography, more sophisticated tools including magnetic resonance imaging, positron emission tomography, and pathological evaluation of biopsies, have identified discrete anatomic abnormalities in many patients diagnosed with the Brugada syndrome. The somewhat ambiguous and complex relationship between clinical phenotype of Brugada syndrome, the presence of SCN5A gene mutations, and structural heart disease has been brought into sharper focus by recent clinical and basic studies.

The suggestion that patients with the clinical features of Brugada syndrome can have underlying structural abnormalities stems back to 1996 when Corrado et al. described a family with autosomal dominant inheritance of clinical features of Brugada syndrome, but in whom postmortem investigation disclosed RV dilation, fibrofatty replacement of the RV free wall, and fibrotic disruption of the right bundle branch. In 2001, Corrado and co-workers reported another study of 273 young sudden death victims from the Veneto region of Italy with the Brugada syndrome phenotype. Twelve-lead ECGs were available from 96 patients and 14% (12 males and 2 females) displayed right precordial ST-segment elevation. At autopsy, 12 of 13 showed pathological features of ARVC, and 1 had an apparently normal heart.

A recent study by Frustaci and co-workers examined 18 consecutive patients (15 males and 3 females) with typical ECG features of Brugada syndrome, who were found to have normal cardiac structure and function based on conventional noninvasive analysis using two-dimensional echocardiography. VF was documented in seven patients, sustained polymorphic VT in seven, and syncope in four. All patients underwent biventricular endomyocardial biopsies obtained 1 to 3 months after the last known arrhythmic event in an effort to limit the possibility that any observed structural abnormalities might have been caused...
directly by the arrhythmia. In addition, the entire coding region of the SCN5A gene was analyzed in all patients; and in some patients, the coding regions of RyR2 (the cardiac ryanodine receptor) and the PKP2 (plakophilin-2) genes were screened. Surprisingly, pathologic features were found in biopsies from all 18 patients included in this study. Most patients had evidence of myocarditis. Diffuse or localized RV inflammation was observed in 14 patients (78%), and viral genomes were detectable by polymerase chain reaction in 4 of them (Coxsackie B3 in 2 patients, and Epstein-Barr virus and Parvovirus B-19 in single patients). Interestingly, the typical ST-segment abnormalities in the right precordial leads that formed the basis of the diagnosis of Brugada syndrome disappeared a few weeks after hospital discharge and were not observed again at follow-up in eight patients in whom endomyocardial biopsy revealed myocardial inflammation. An SCN5A mutation was not identified in these patients. These observations serve to confirm the fact that myocarditis is capable of producing an acquired form of the Brugada syndrome.48 Other types of cardiac muscle pathology were observed in biopsies from the remaining four patients. One patient showed extensive fibrofatty replacement typical of ARVC in the RV biopsy and the other three nonspecific abnormalities consistent with diffuse muscle disease typically present in dilated cardiomyopathy. Overall, RV biopsy was abnormal in all 18 patients, whereas the left ventricle biopsy was considered normal in 10 patients.

Novel SCN5A mutations associated with loss of function were detected in 4 of the 18 patients (22%). Interestingly, none had myocarditis on their biopsies, although three of the four exhibited myocyte hypertrophy and vacuolization, changes that should be viewed with some caution in that they are rather nonspecific, given the usual amount of artifactual tissue disruption that occurs during the endomyocardial biopsy procedure.49

Although the study by Frustaci and co-workers47 is not conclusive, it is consistent with the notion that primary channelopathies can lead to cellular changes.50 Age-related development of myocardial fibrosis and progressive slowing of atrial and ventricular conduction have been reported in mice with a single null allele of the SCN5A gene (SCN5A+/- mice).51

That SCN5A mutations can be associated with structural cardiac defects in humans was reported by Bezzina and co-workers in 2003.52 This was a family whose members exhibited irregular wide-complex tachycardia and in whom mutational analysis revealed compound heterozygous mutations in SCN5A. Pathological analysis of the heart in one family member revealed changes of dilated cardiomyopathy with severe degenerative abnormalities of the cardiac conduction system.

Frustaci et al.47 suggested that deranged intracellular sodium homeostasis caused by a SCN5A mutation could act via sodium-hydrogen and sodium-calcium exchangers to affect intracellular pH and calcium flux, which, in turn, could impair excitation-contraction coupling and energy production. Another hypothesis is that loss of the AP, which is thought to provide the arrhythmogenic substrate in Brugada syndrome, leads to a hibernation-like state that is associated with progressive cellular changes.50,53 The all-or-none repolarization at the end of phase 1 of the AP responsible for loss of the dome causes the calcium channel to inactivate very soon after it activates. As a consequence, calcium channel current is drastically reduced, the cell becomes depleted of calcium within seconds, and contractile function ceases. This leads to wall motion abnormalities, observed in many patients with Brugada syndrome. Loss of the AP dome, because it creates a hibernation-like state, may over long periods of time lead to mild structural changes, including intracellular lipid accumulation and vacuolization. These structural changes may in turn contribute to the arrhythmogenic substrate of the Brugada syndrome, although they are very different from those encountered in ARVC/D.50,53

Whatever the mechanism, structural remodeling of the myocardium in a patient with an SCN5A mutation could help to explain the delay in manifestation of ECG abnormalities and arrhythmias, and may account for the eventual progression of some cases of Brugada syndrome to ones exhibiting structural abnormalities.

These studies also point to inflammatory processes as being primary and in some cases implicating myocarditis in the development of an acquired form of Brugada syndrome, whose manifestation may be transient.47 To what extent the electrocardiographic and arrhythmic manifestations of primary structural defects may have a genetic predisposition is not known and must await delineation of the 70–80% of Brugada syndrome cases that have thus far not been linked to a genetic mutation.

These facts notwithstanding, the majority of congenital Brugada syndrome patients are believed to possess a structurally normal heart, consistent with the notion that this is a primary electrical heart disease.54 While fibrosis and myocarditis may exacerbate or indeed trigger events in patients with the Brugada syndrome, it seems clear that in the vast majority of cases these structural changes are unrelated to ARVC/D. ARVC and Brugada syndromes are distinct clinical entities, both with respect to the clinical presentation and...
The only genes thus far linked to the Brugada syndrome are those that influence sodium channel current, whereas ARVC has been linked to nine different chromosomal loci and four putative genes (desmoplakin, plakoglobin, RyR2, and plakophilin-2) independent of those responsible for the Brugada syndrome. Only the ARVC5 locus has been mapped to a region overlapping with the second locus for Brugada syndrome, but the responsible gene has not been identified. In Brugada syndrome, imaging techniques such as echocardiography, angiography, magnetic resonance imaging, and radionuclide scintigraphy show no evidence of overt structural heart disease, whereas ARVC patients characteristically display RV morphological and functional changes (such as global diastasis, bulgings/aneurysms, and wall motion abnormalities). Ventricular arrhythmias in ARVC are most commonly monomorphic VT, often precipitated by catecholamines or exercise, accounting for sudden death of young competitive athletes. In contrast, ST-segment elevation and arrhythmias in Brugada patients are enhanced by vagotonic agents or $\beta$-adrenergic blockers, and polymorphic VT most commonly occurs during rest or sleep. Unlike Brugada syndrome, the ECG abnormalities in ARVC are usually not dynamic, displaying a constant T-wave inversion, epsilon waves and, in the progressive stage, reduction of the R amplitude, which are largely unaffected by sodium channel blocker administration.

Recent studies report that electron beam computed tomography has uncovered wall motion abnormalities in a series of Brugada patients tested. Although such contractile abnormalities are commonly considered pathognomonic of structural disease, recent studies suggest that such contractile dysfunction can result from loss of the AP dome in regions of RV epicardium, and, thus, may be unrelated to any type of morphological defect. Loss of the dome leads to contractile dysactivity of non-crest neuron-epithelial cells that mediate the development of the neural crest, 66 and its absence is known to result in aberrant cardiac infundibulum and neural crest formation. Germline knockout (KO) of Cx43 in the mouse results in abnormal cardiac morphogenesis, including RVOT abnormalities and perinatal death. Liu et al. recently tested the hypothesis that Cx43 expression in the cardiac neural crest is critical to normal development of the outflow tracts using a CKO strategy to generate mice with neural-crest-restricted KO of Cx43. The CKO mice displayed marked dilatation or bulging of the outflow tracts. The results suggested that Cx43 expression is not required in the neural crest or dorsal neural tube for normal outflow tract morphogenesis, but is critically important in the remainder of the neural tube, where it may regulate the plasticity of non-crest neuron-epithelial cells that mediate the development of the outflow tracts. It is tempting to speculate that Cx43 defects limited to this region of the heart may contribute to the development of the Brugada syndrome phenotype by impairing electrotonic communication in the RVOT. This would exacerbate repolarization heterogeneities which contribute prominently to the arrhythmogenic substrate, as well as to slowing of conduction, which may facilitate the development of reentry.

Risk Stratification

A great deal of focus and debate has centered around the issue of risk stratification of patients at risk for sudden cardiac death. It is generally accepted that Brugada syndrome patients presenting with aborted sudden death are at high risk for
Brugada et al.71 reported that only 6% of asymptomatic patients had become symptomatic; an occurrence reported that 8% of previously asymptomatic patients displayed a first event during a follow-up period of 27±29 months, the same authors69 reported that 8% of previously asymptomatic patients had become symptomatic; an occurrence of a life-threatening event of 3.5%/year. In 2005, the same authors reported that 0.35%/year. In 2005, the same authors reported that 1 out of 123 asymptomatic individuals with a Type 1 ECG (0.8%) had a first arrhythmic event during a 40 ± 50 month follow-up. This translates into a first event rate of 0.24% per year, considerably less than the other two registries.

The major registry studies all agree that Brugada syndrome patients at higher risk for the development of subsequent events are those presenting with a spontaneous Type 1 ST-segment elevation or Brugada ECG and/or those with previous VT/VF or SCD.73 The registries also agree that PES inducibility is greatest among patients with previous VT/VF or syncope. Approximately one-third of asymptomatic patients are inducible. In the Priori and Napolitano72 and Eckardt et al.73 studies, inducibility of VT/VF in asymptomatic patients was not associated with risk. The lack of association between inducibility and spontaneous VF in Brugada patients was also reported by a number of smaller studies, such as that of Kanda et al.77 In sharp contrast, Brugada et al.78 found that the risk for developing VT/VF is much greater in patients who were inducible during PES, whether or not a Type 1 ST-segment elevation was spontaneously present and whether or not they were symptomatic. The reason for the marked disparity in the predictive power of PES inducibility among the different studies is not immediately apparent. The discrepancies may be due to differences in patient characteristics and the use of multiple testing centers with non-standardized or non-comparable stimulation protocols.79 Additional studies are needed to further define risk stratification strategies for asymptomatic patients.

It is noteworthy that in experimental models of the Brugada syndrome involving the coronary-perfused wedge preparation, polymorphic VT is readily inducible with a single S2, but only when applied on the epicardial surface of the wedge. Inducibility is not possible or much more difficult when extra-stimulation is applied to the endocardial surface. The shorter refractory period of epicardial preparations may contribute to the discrepancy in the incidence of first events among asymptomatic patients reported by the two largest registries of Brugada syndrome patients, a recent report by Eckardt and co-workers74 suggests that other factors may be involved. These authors report that 1 out of 123 asymptomatic individuals with a Type 1 ECG (0.8%) had a first arrhythmic event during a 40 ± 50 month follow-up. This translates into a first event rate of 0.24% per year, considerably less than the other two registries.

The reason behind the disparity in the data generated by these two groups is not clearly the case, it may be a rare occurrence, and the exclusion appears justified on the basis that it avoids a large number of false positive diagnoses. The failure to exclude individuals with Type 2 and 3 ST-segment elevation suggests that the European registry may contain many individuals who do not have the syndrome.

While it stands to reason that this difference may contribute to the discrepancy in the incidence of first events among asymptomatic patients, the reason for the marked disparity in the predictive power of PES inducibility among the different studies is not immediately apparent. The discrepancies may be due to differences in patient characteristics and the use of multiple testing centers with non-standardized or non-comparable stimulation protocols. Additional studies are needed to further define risk stratification strategies for asymptomatic patients.

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thus facilitating the induction of reentry. These relationships suggest that PES applied to the epicardium may provide a more accurate assessment of risk than the current approach in which stimuli are applied to the endocardial surface. In support of this hypothesis, Carlsson et al. reported that a Brugada syndrome patient with recurrent syncope due to polymorphic VT could not be induced with RV endocardial stimulation. However, epicardial stimulation from a left ventricular site through the coronary sinus led to the development of polymorphic VT.80

Recent studies have suggested that combined electrocardiographic markers may be helpful in risk stratification. Atarashi and Ogawa74 used the electrocardiographic markers may be helpful in risk stratification.80

Hevia and co-workers28 recently evaluated Tp-e as a risk factor for recurrence of life-threatening cardiac events in patients with the Brugada syndrome. Twenty-nine patients with the ECG pattern of Brugada syndrome and 29 healthy age- and sex-matched controls were studied. QT, QTc, QT dispersion, Tp-e, and Tp-e dispersion were measured. Eleven patients with the ECG pattern of Brugada syndrome had a prolonged (>460 ms) QTc in V2, but usually not in inferior or left leads. No patient had abnormally prolonged QT dispersion. Tp-e and Tp-e dispersion were significantly prolonged in patients with recurrences versus patients without events. Thus, there was a significant correlation between Tp-e and Tp-e dispersion and occurrence of life-threatening arrhythmic events in patients with the Brugada syndrome, suggesting that these parameters may be useful in risk stratification of patients with the syndrome.

Genetic Factors Underlying the Brugada Syndrome

Brugada syndrome is inherited via an autosomal dominant mode of transmission. The first gene to be linked to the Brugada syndrome is SCN5A, the gene encoding for the α-subunit of the cardiac sodium channel gene.90 Figure 3 highlights the diversity of SCN5A mutations associated with the Brugada syndrome. Of note, mutations in SCN5A are also responsible for the LQT3 form of the long-QT syndrome and cardiac conduction disease. A number of mutations have been reported to cause overlapping syndromes; in some cases, all three phenotypes are present.91

Over one hundred mutations in SCN5A have been linked to the syndrome in recent years (see Antzelevitch et al.61 for references; also see www.fsm.it/cardmoc). Only a fraction of these mutations have been studied in expression systems and shown to result in loss of function due either to: (1) failure of the sodium channel to express; (2) a shift in the voltage- and time-dependence of sodium channel current (INa) activation, inactivation, or reactivation; (3) entry of the sodium channel into an intermediate state of inactivation from which it recovers more slowly; or (4) accelerated inactivation of the sodium channel. In vitro expression systems, the premature inactivation of the sodium channel is sometimes observed at physiological temperatures, but not at room
Temperature. Acceleration of $I_{Na}$ inactivation was still more accentuated at higher than physiological temperatures, suggesting that the syndrome may be unmasked, and that patients with the Brugada syndrome may be at an increased risk, during a febrile state. A number of Brugada patients displaying fever-induced polymorphic VT have been identified since the publication of this report.

Mutations in the $SCN5A$ gene account for approximately 18–30% of Brugada syndrome cases. A higher incidence of $SCN5A$ mutations has been reported in familial than in sporadic cases. Of note, negative $SCN5A$ results generally do not rule out causal gene mutations, since the promoter region, cryptic splicing mutations, or presence of gross rearrangements are generally not part of routine investigation. A recent report by Hong et al. provided the first report of a dysfunctional sodium channel created by an intronic mutation giving rise to cryptic splice site activation in $SCN5A$ in a family with the Brugada syndrome. The deletion of fragments of segments 2 and 3 of Domain IV of $SCN5A$ caused complete loss of function.

Bezzina and co-workers recently provided interesting evidence in support of the hypothesis that an $SCN5A$ promoter polymorphism common in Asians modulates variability in cardiac conduction, and may contribute to the high prevalence of Brugada syndrome in the Asian population. Sequencing of the $SCN5A$ promoter identified a haplotype variant consisting of 6 polymorphisms in near-complete linkage disequilibrium that occurred at an allele frequency of 22% in Asian subjects and was absent in whites and blacks. The results of the study demonstrate that sodium channel transcription in the human heart may vary considerably among individuals and races, and be associated with variable conduction velocity and arrhythmia susceptibility.

A second locus on chromosome 3, close to but distinct from $SCN5A$, has been linked to the syndrome in a large pedigree in which the syndrome is associated with progressive conduction disease, a low sensitivity to procainamide, and a relatively good prognosis. The gene was recently identified as the Glycerol-3-Phosphate Dehydrogenase 1-Like Gene ($GPD1L$). A mutation in $GPD1L$ has been shown to result in a partial reduction of $I_{Na}$.

Knowledge thus far gained through genetic analysis suggests that identification of specific mutations may not be very helpful in formulating a diagnosis or providing a prognosis. There are no clear hotspots and mutations have been reported throughout the $SCN5A$ gene. It is not clear whether some mutations are associated with a greater risk of arrhythmic events or sudden death. Genetic testing is recommended for support of the clinical diagnosis, for early detection of relatives at potential risk, and particularly for the purpose of advancing research and consequently our understanding of genotype-phenotype relations.

Cellular and Ionic Mechanisms Underlying the Development of the Brugada Phenotype

The concept of phase 2 reentry, which is a known trigger for the Brugada syndrome, was described in the early 1990s and evolved in parallel with the clinical discovery of the Brugada syndrome. Studies conducted over the past decade suggest that rebalancing of the currents active at the end of phase 1, leading to an
Figure 4. Phase 2 reentry. Reentrant activity induced by exposure of a canine ventricular epicardial preparation (0.7 cm²) to simulated ischemia. Microelectrode recordings were obtained from four sites as shown in the schematic (upper right). After 35 min of ischemia, the action potential dome develops normally at site 4, but not at sites 1, 2, or 3. The dome then propagates in a clockwise direction re-exciting sites 3, 2, and 1 with progressive delays, thus generating a closely coupled reentrant extrasystole (156 ms) at site 1. In this example of phase 2 reentry, propagation of the dome occurs in a direction opposite to that of phase 0, a mechanism akin to reflection. BCL = 700 ms. Modified from Lukas and Antzelevitch.119 with permission.

accentuation of the AP notch in RV epicardium is responsible for the accentuated J wave or ST-segment elevation associated with the Brugada syndrome.105-115 Under normal conditions, the appearance of the epicardial AP notch is due principally to the interaction of two ion channel currents. The transient outward current (I_{to}), which activates during phase 0, contributes most promi-

nently to phase 1 of the AP, whereas the calcium inward current is largely responsible for the second upstroke, giving rise to the AP dome. Amplification of epicardial and TDR secondary to the presence of genetic defects, pathophysiologic factors, and pharmacologic influences, leads to accentuation of the J wave and eventually to loss of the AP dome, giving rise to extrasystolic activity in the form of phase 2 reentry. Activation of I_{to} leads to a paradoxical prolongation of APD in canine ventricular tissues,116 but to abbreviation of ventricular APD in species that normally exhibit brief APs (e.g., mouse and rat).117 Pathophysiologic conditions (e.g., ischemia, metabolic inhibition) and some pharmacologic interventions (e.g., I_{Na} or I_{Ca} blockers or I_{K-ATP}, I_{to}, I_{Ks}, or I_{Ks} activators) can lead to marked abbreviation of the AP in canine and feline118 ventricular cells where I_{to} is prominent. Under these conditions, canine ventricular epicardium exhibits an all-or-none repolarization as a result of the shift in the balance of currents flowing at the end of phase 1 of the AP. When phase 1 reaches approximately −30 mV, all-or-none repolarization of the AP ensues, leading to loss of the dome as the outward currents overwhelm the inward currents. Loss of the AP dome generally occurs at some epicardial sites but not others, resulting in the development of a marked dispersion of repolarization within the epicardium as well as transmurally, between epicardium and endocardium. Propagation of the AP dome from the epicardial site at which it is maintained to sites at which it is abolished can cause local re-excitation of the preparation. This mechanism, termed phase 2 reentry, produces extrasystolic beats capable of initiating circus movement reentry119 (Fig. 4). Phase 2 reentry has been shown to occur when RV epicardium is exposed to: (1) K⁺ channel openers such as pinacidil120; (2) sodium channel blockers such as flecainide121; (3) increased [Ca²⁺]o122; (4) calcium channel blockers such as verapamil; (5) metabolic inhibition123; and (6) simulated ischemia.119

Exaggerated or otherwise abnormal J waves have long been linked to idiopathic VF and the Brugada syndrome.1,19,42,112,124,125 The Brugada syndrome is characterized by exaggerated J wave that manifests as an ST-segment elevation in the right precordial leads.1 A number of studies have highlighted the similarities between the conditions that predispose to phase 2 reentry and those that attend the appearance of the Brugada syndrome. Loss of the AP dome in epicardium, but not endocardium, generates a transmural current that manifests on the ECG as an ST-segment elevation, similar to that encountered in patients with the Brugada syndrome.105,123,126 Evidence in
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Figure 5. ECG and arrhythmias with typical features of the Brugada syndrome recorded from canine right ventricular wedge preparations. (A) Schematic of arterially perfused right ventricular wedge preparation. (B) Pressure-induced phase 2 reentry and VT. Shown are transmembrane action potentials simultaneously recorded from two epicardial (Epi 1 and Epi 2) and one M region (M) sites, together with a transmural ECG. Local application of pressure near Epi 2 results in loss of the action potential dome at that site but not at Epi 1 or M sites. The dome at Epi 1 then re-excites Epi 2, giving rise to a phase 2 reentrant extrasystole which triggers a short run of ventricular tachycardia. Note the ST-segment elevation due to loss of the action potential dome in a segment of epicardium. (C) Polymorphic VT/VF induced by local application of the potassium channel opener pinacidil (10 μM) to the epicardial surface of the wedge. Action potentials from two epicardial sites (Epi 1 and Epi 2) and a transmural ECG were simultaneously recorded. Loss of the dome at Epi 1 but not Epi 2 creates a marked dispersion of repolarization, giving rise to a phase 2 reentrant extrasystole. The extrasystolic beat then triggers a long episode of ventricular fibrillation (22 sec). (Right panel) Addition of 4-aminopyridine (4-AP, 2 mM), a specific Ito blocker, to the perfusate restored the action potential dome at Epi 1, thus reducing dispersion of repolarization and suppressing all arrhythmic activity. BCL = 2,000 ms. (D) Phase 2 reentry gives rise to VT following addition of pinacidil (2.5 μM) to the coronary perfusate. Transmembrane action potentials form 2 epicardial sites (Epi 1 and Epi 2) and one endocardial site (Endo), as well as a transmural ECG were simultaneously recorded. (Right panel) 4-AP (1 mM) markedly reduces the magnitude of the action potential notch in epicardium, thus restoring the action potential dome throughout the preparation and abolishing all arrhythmic activity. Panel D is from Yan and Antzelevitch106 with permission.

support of a phase 2 reentrant mechanism in humans was recently provided by Thomsen et al.127 and Antzelevitch.128

Autonomic neurotransmitters like acetylcholine facilitate loss of the AP dome129 by suppressing I_{Ca} and/or augmenting potassium current. β-adrenergic agonists restore the dome by augmenting I_{Ca}. Sodium channel blockers also facilitate loss of the canine RV AP dome via a negative shift in the voltage at which phase 1 begins.121,130 These findings are consistent with accentuation of ST-segment elevation in patients with the Brugada syndrome following vagal maneuvers or Class I antiarrhythmic agents, as well as normalization of the ST-segment elevation following β-adrenergic agents and phosphodiesterase III inhibitors.13,105,131 Loss of the AP dome is much more readily induced in right versus left canine ventricular epicardium123,126,132 because of the more prominent I_{to}-mediated phase 1 in APs in
Figure 6. Terfenadine-induced ST-segment elevation, T-wave inversion, transmural and endocardial dispersion of repolarization, and phase 2 reentry. Each panel shows transmembrane action potentials from one endocardial (top) and two epicardial sites together with a transmural ECG recorded from a canine arterially perfused right ventricular wedge preparation. (A) Control (BCL 400 ms). (B) Terfenadine (5 $\mu$M) accentuated the epicardial action potential notch creating a transmural voltage gradient that manifests as a ST-segment elevation or exaggerated J wave in the ECG. First beat recorded after changing from BCL 800 ms to BCL 400 ms. (C) Continued pacing at BCL 400 ms results in all-or-none repolarization at the end of phase 1 at some epicardial sites but not others, creating a local epicardial dispersion of repolarization (EDR) as well as a transmural dispersion of repolarization (TDR). (D) Phase 2 reentry occurs when the epicardial action potential dome propagates from a site where it is maintained to regions where it has been lost. Modified from Fish and Antzelevitch$^{107}$ with permission.

This region of the heart. This distinction is believed to be the basis for why the Brugada syndrome is a RV disease.

Thus, accentuation of the RV epicardial AP notch underlies the ST-segment elevation. Eventual loss of the dome of the RV epicardial AP further exaggerates ST-segment elevation. The vulnerable window created within the epicardium, as well as transmurally, serves as the substrate and phase 2 reentry provides the extrasystole that serves as the trigger which precipitates episodes of VT and VF in the Brugada syndrome. Evidence in support of this hypothesis was recently provided in an arterially perfused canine RV experimental model of the Brugada syndrome (Fig. 5).$^{106}$ The VT and VF generated in these preparations is usually polymorphic, resembling a rapid form of TdP. This activity may be mechanistically related to the migrating spiral wave shown to generate a pattern resembling TdP associated with a normal or long-QT interval.$^{133,134}$

Much of the focus in the past has been on the ability of a reduction in sodium channel current to unmask the Brugada syndrome and create an arrhythmogenic substrate. A recent report shows that a combination of $I_{Na}$ and $I_{Ca}$ block is more effective than $I_{Na}$ inhibition alone in precipitating the Brugada syndrome in the arterially perfused wedge preparation (Fig. 6).$^{107}$ High concentrations of terfenadine (5 $\mu$M) produce a potent block of $I_{Na}$ and $I_{Ca}$, leading to accentuation of the epicardial AP notch following acceleration of the rate from a basic cycle length of 800 to 400 ms. The dramatic accentuation of the notch was due to the effect of the drug to depress phase 0, augment the magnitude of phase 1, and delay the appearance of the second upstroke. With continued rapid pacing, phase 1 becomes more accentuated, until all-or-none repolarization occurs at the end of phase 1 at some epicardial sites but not others, leading to the development of both epicardial dispersion of repolarization (EDR) and TDR (Fig. 6C). Propagation of the dome from the region where it is maintained to the region at which it is lost results in the development of local phase 2 reentry (Fig. 6D). Figure 7 shows the ability of terfenadine-induced phase 2 reentry to generate an extrasystole, couplet, and polymorphic VT/VF. Figure 7D illustrates an
example of PES to initiate VT/VF under similar conditions.

The electrocardiographic manifestations of the Brugada syndrome have been attributed to: (1) conduction delay in the RV epicardial free wall in the region of the outflow tract (RVOT)\textsuperscript{135} and/or (2) premature repolarization of the RV epicardial AP secondary to loss of the AP dome, or a combination of the two.\textsuperscript{136} The cellular mechanisms thought to be responsible for the development of the Brugada phenotype via hypothesis 2 is schematically illustrated in Figure 8.\textsuperscript{137,138}

Under physiological conditions, the ST segment is isoelectric because of the absence of transmural voltage gradients at the level of the AP plateau (Fig. 8A). Accentuation of the RV notch under pathophysiologic conditions leads to exaggeration of transmural voltage gradients and thus to accentuation of the J wave or to J point elevation. When epicardial repolarization precedes repolarization of the cells in the M and endocardial regions, the T wave remains positive. The result is a saddleback configuration of the repolarization waves (Fig. 8B). Further accentuation of the notch may be accompanied by a prolongation of the epicardial AP such that the direction of repolarization across the RV wall and transmural voltage gradients are reversed, leading to the development of a coved-type ST-segment elevation and inversion of the T wave (Fig. 8C), typically observed in the ECG of Brugada patients. A delay in epicardial activation may also contribute to inversion of the T wave. The downsloping ST-segment elevation observed in the experimental wedge models often appears as an R’, suggesting that the appearance of a right bundle branch block (RBBB) morphology in Brugada patients may be due at least in part to early repolarization of RV epicardium, rather than major impulse conduction block in the right bundle.

A rigorous application of RBBB criteria reveals that a large majority of RBBB-like morphologies encountered in cases of Brugada syndrome do not fit the criteria for RBBB.\textsuperscript{139} Moreover, attempts by Miyazaki and co-workers to record delayed activation of the RV in Brugada patients met with failure.\textsuperscript{13}

It is important to point out that although the typical Brugada morphology is present in Figure 8B,C, an arrhythmogenic substrate is
absent. The arrhythmogenic substrate is thought to develop when a further shift in the balance of current leads to loss of the AP dome at some epicardial sites but not others (Fig. 8D). Loss of the AP dome in epicardium but not endocardium results in the development of a marked TDR and refractoriness, responsible for the development of a vulnerable window during which a premature impulse or extrasystole can induce a reentrant arrhythmia. Conduction of the AP dome from sites at which it is maintained to sites at which it is lost causes local re-excitation via a phase 2 reentry mechanism, leading to the development of a very closely coupled extrasystole, which captures the vulnerable window across the wall, thus triggering a circus movement reentry in the form of VT/VF (Fig. 8E). A high resolution optical mapping system that allows simultaneous recording of transmembrane APs from 256 sites along the transmural surface of the arterially perfused canine RV wedge preparation has been used by Shimizu et al. to demonstrate that a steep repolarization gradient between the region at which the dome is lost and the region at which it is maintained is essential for the development of a closely coupled phase 2 reentrant extrasystole. This study also showed that reentry initially rotates in the epicardium and gradually shifts to a transmural orientation, responsible for nonsustained polymorphic VT or VF.

A recent report by Kurita et al., in which monophasic AP electrodes were positioned on the epicardial and endocardial surfaces of the RVOT in patients with the Brugada syndrome, provides further support. The phase 2 reentrant beat fuses with the negative T wave of the basic response. Because the extrasystole originates in epicardium, the QRS complex is largely comprised of a Q wave, which serves to accentuate the negative deflection of the inverted T wave, giving the ECG a more symmetrical appearance. This morphology is often observed in the clinic preceding the onset of polymorphic VT.

Studies involving the arterially perfused RV wedge preparation provide evidence in support of these hypotheses. A high resolution optical mapping system that allows simultaneous recording of transmembrane APs from 256 sites along the transmural surface of the arterially perfused canine RV wedge preparation has been used by Shimizu et al. to demonstrate that a steep repolarization gradient between the region at which the dome is lost and the region at which it is maintained is essential for the development of a closely coupled phase 2 reentrant extrasystole. This study also showed that reentry initially rotates in the epicardium and gradually shifts to a transmural orientation, responsible for nonsustained polymorphic VT or VF.

The marked accentuation of the epicardial AP dome and the development of concealed phase 2 reentry suggest that activation forces may extend beyond the QRS in Brugada patients. Indeed, signal averaged ECG (SAECG) recordings have demonstrated late potentials in patients with the Brugada syndrome, especially in the anterior wall of the RVOT. The basis for these late

**Figure 8.** Schematic representation of right ventricular epicardial action potential changes proposed to underlie the electrocardiographic manifestation of the Brugada syndrome. Modified from Antzelevitch with permission.
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potentials, commonly ascribed to delayed conduction within the ventricle, is largely unknown. Endocardial recordings have been unrevealing. Nagele and co-workers\textsuperscript{143} introduced a guide wire into the conus branch of the right coronary artery to record signals from the epicardial surface of the anterior wall of the RVOT in patients with the Brugada syndrome. The unipolar recordings displayed delayed potentials, which coincided with late potentials recorded in the SAECG, particularly after administration of Class IC antiarrhythmic agents. The authors conclude that recordings from the conus branch of the right coronary artery can identify an “epicardial abnormality” in the RVOT, which is accentuated in the presence of IC agents, thus uncovering part of the arrhythmogenic substrate responsible for VT/VF in Brugada syndrome, which may be related to the second upstroke or a concealed phase 2 reentrant beat. Late potentials are often regarded as being representative of delayed activation of the myocardium, but in the case of the Brugada syndrome, other possibilities exist as discussed above. For example, the second upstroke of the epicardial AP, thought to be greatly accentuated in Brugada syndrome,\textsuperscript{137} might be capable of generating late potentials when RVOT activation is otherwise normal. Moreover, the occurrence of phase 2 reentry, especially when concealed (i.e., when it fails to trigger transmural reentry), may contribute to the generation of delayed unipolar and late SAECG potentials.

The rate-dependence of the ECG sign can be used to discriminate between these two hypotheses. If the Brugada ECG sign is due to delayed conduction in the RVOT, acceleration of the rate would be expected to further aggravate conduction and thus accentuate the ST-segment elevation and the RBBB morphology of the ECG. If, on the other hand, the Brugada ECG sign is secondary to accentuation of the epicardial AP notch, at some point leading to loss of the AP dome, acceleration of the rate would be expected to normalize the ECG, by restoring the AP dome and reducing the notch. This occurs because the transient outward current, which is at the heart of this mechanism, is slow to recover from inactivation and is less available at faster rates. The fact of the matter is that Brugada patients usually display a normalization of their ECG or no change when heart rate is increased, thus favoring the second hypothesis. Further evidence in support of this hypothesis derives from the recent observations of Shimizu and co-workers\textsuperscript{146} that quinidine suppresses late potentials recorded in a patient with Brugada syndrome. This effect of the drug is presumably due to inhibition of I\textsubscript{to}, leading to diminution of the epicardial AP notch and normalization of the repolarization heterogeneities. If the late potentials were due to delayed conduction, quinidine-induced I\textsubscript{Na} inhibition would be expected to accentuate the appearance of the late potentials.

Thus, the available data, both basic and clinical, point to transmural voltage gradients that develop secondary to accentuation of the epicardial notch and loss of the AP dome as being in large part responsible for the Brugada ECG signature.

Gender-Based Differences in the Manifestation of Brugada Syndrome

Although the genetic mutation responsible for the Brugada syndrome is equally distributed between the sexes, the clinical phenotype is 8 to 10 times more prevalent in males than in females. The basis for this sex-related distinction has been shown to be due to a more prominent I\textsubscript{to}-mediated AP notch in the RV epicardium of males versus females\textsuperscript{148} (Figs. 9 and 10). The more prominent I\textsubscript{to} causes the end of phase 1 of the RV epicardial AP to repolarize to more negative potentials in tissue and arterially perfused wedge preparations from males, facilitating loss of the AP dome and the development of phase 2 reentry and polymorphic VT.

The proposed cellular mechanism for the Brugada syndrome is summarized in Figure 11. The available data support the hypothesis that the Brugada syndrome results from amplification of heterogeneities intrinsic to the early phases of the AP among the different transmural cell types. The amplification is secondary to a rebalancing of currents active during phase 1, including a decrease in I\textsubscript{Na} or I\textsubscript{Ca}, or augmentation of any one of a number of outward currents. ST-segment elevation similar to that observed in patients with the Brugada syndrome occurs as a consequence of the accentuation of the AP notch, eventually leading to loss of the AP dome in RV epicardium, where I\textsubscript{Na} is most prominent. Loss of the dome gives rise to both a TDR as well as EDR. The transmural dispersion is responsible for the development of ST-segment elevation and the creation of a vulnerable window across the ventricular wall, whereas the epicardial dispersion gives to phase 2 reentry which provides the extrasystole that captures the vulnerable window, thus precipitating VT/VF. The
VT generated is usually polymorphic, resembling a very rapid form of TdP.

Modulating Factors and Acquired Forms of Brugada Syndrome

A number of factors modulate the electrocardiographic and arrhythmic manifestations of the Brugada syndrome. ST-segment elevation in the Brugada syndrome is often dynamic. The Brugada ECG may often be concealed, but can be unmasked or modulated by sodium channel blockers, a febrile state, vagotonic agents, \( \alpha \)-adrenergic agonists, \( \beta \)-adrenergic blockers, tricyclic or tetracyclic antidepressants, first generation antihistaminics (dimenhydrinate), a combination of glucose and insulin, hyperkalemia, hypokalemia, hypercalcemia, and by alcohol and cocaine toxicity (Fig. 12).11–13,149–156 These agents may also induce acquired forms of the Brugada syndrome (Table IV). Although a definitive list of drugs to avoid in the Brugada syndrome has not yet been formulated, the list of agents in Table IV may provide some guidance. One of the more recent additions to this group is Lithium. This widely used drug is a potent blocker of cardiac sodium channels and can unmask patients with the Brugada syndrome.157

Myocardial infarction or acute ischemia due to vasospasm involving the RVOT mimics ST-segment elevation similar to that in Brugada syndrome. This effect is secondary to the depression of \( I_{Ca} \) and the activation of \( I_{KATP} \) during ischemia, and suggests that patients with congenital and possibly acquired forms of Brugada syndrome may be at a higher risk for ischemia-related sudden cardiac death.158 Although the coexistence of Brugada syndrome and vasospastic angina in the same patient is not rare, Chinushi et al. have failed to observe an enhanced susceptibility to VF nor a proarrhythmic effect of Ca-antagonist in this setting.159

VT/VF and sudden death in the Brugada syndrome usually occur at rest and at night. Circadian variation of sympatho-vagal balance, hormones, and other metabolic factors are likely to contribute to this circadian pattern. Bradycardia, due to altered sympatho-vagal balance or other factors, may contribute to arrhythmia initiation.32,160,161 Abnormal \( ^{123}\text{I-MIBG} \) uptake in 8 (17%) of the 17 Brugada syndrome patients but none in the control group was demonstrated by Wichter et al.162 There was segmental reduction of \( ^{123}\text{I-MIBG} \) in the inferior and the septal left ventricular wall, indicating presynaptic sympathetic dysfunction. Of note, imaging of the right ventricle, particularly the RVOT, is difficult with this technique, so that insufficient information is available concerning sympathetic function in the regions known to harbor the arrhythmogenic substrate. Moreover, it remains unclear what role the reduced uptake function plays in the arrhythmogenesis of the Brugada syndrome. If indeed the RVOT is similarly affected, this defect may indeed alter the sympathovagal balance in favor of the development of an arrhythmogenic substrate.106,129

Hypokalemia has been implicated as a contributing cause for the high prevalence of SUDS in the northeastern region of Thailand where potassium deficiency is endemic.164,155 Serum potassium in the northeastern population is significantly lower than that of the population in Bangkok, which lies in the central part of Thailand where potassium is abundant in the food.
A recent case report highlights the ability of hypokalemia to induce VF in a 60-year-old man who had asymptomatic Brugada syndrome, without a family history of sudden cardiac death. This patient was initially treated for asthma by steroids, which lowered serum potassium from 3.8 mmol/L on admission to 3.4 and 2.9 mmol/L on the 7th day and 8th day of admission, respectively. Both were associated with unconsciousness. VF was documented during the last episode, which reverted spontaneously to sinus rhythm.

The Thai Ministry of Public Health Report (1990) found an association between a large meal of glutinous rice (“sticky rice”) or carbohydrates ingested on the night of death in SUNDS patients. Consistent with this observation, a recent study by Nogami et al. found that glucose and insulin could unmask the Brugada ECG. Another possibility is that sudden death in these patients is due to the increased vagal tone produced by the stomach distention. A recent study by Ikeda et al. has shown that a full stomach after a large meal can unmask a Type I ECG, particularly...
Figure 11. Proposed mechanism for the Brugada syndrome. A shift in the balance of currents serves to amplify existing heterogeneities by causing loss of the action potential dome at some epicardial, but not endocardial, sites. A vulnerable window develops as a result of the dispersion of repolarization and refractoriness within epicardium, as well as across the wall. Epicardial dispersion leads to the development of phase 2 reentry, which provides the extrasystole that captures the vulnerable window and initiates VT/VF via a circus movement reentry mechanism. Modified from Antzelevitch with permission.

Figure 12. Factors predisposing to the electrocardiographic and arrhythmic manifestations of the Brugada syndrome. Modified from Nademanee et al. with permission.
in Brugada syndrome patients at high risks for arrhythmic events, thus suggesting that this technique may be of diagnostic and prognostic value.

Premature inactivation of the sodium channel in SCN5A mutations associated with the Brugada syndrome has been shown to be accentuated at higher temperatures,

92 suggesting that a febrile state may unmask the Brugada syndrome. Indeed, several case reports have emerged recently, demonstrating that febrile illness could reveal the Brugada ECG and precipitate VF.14,93,94,165–167 A recent report from Keller et al. 168 has identified a missense mutation, F1344S, in SCN5A in a patient with Brugada syndrome and fever-induced VT/VF. Expression of F1344S showed a shift in the voltage-dependence of activation, which was further accentuated at high temperatures mimicking fever. Thus fever may also cause a loss of function in INa by producing a shift in the voltage-dependence of activation.

Anecdotal data point to hot baths as a possible precipitating factor. Of note, the Northeastern part of Thailand, where the Brugada syndrome is most prevalent, is known for its very hot climate. A study is underway to assess whether this extreme climate influences the prognosis of the disease.

**Approach to Therapy**

**Device Therapy**

An ICD is the only proven effective device treatment for the disease (Table V).169,170 Recommendations for ICD implantation5,6 are presented in Table VI (from reference 5 with permission) and summarized as follows:

1. **Symptomatic patients** displaying the Type 1 ST-segment elevation or Brugada ECG (either spontaneously or after sodium channel blockade) who present with aborted sudden death should receive an ICD as a Class I indication without additional need for EPS. Similar patients presenting with related symptoms such as syncope, seizure, or nocturnal agonal respiration should also undergo ICD implantation as a Class I indication after non-cardiac causes of these symptoms have been carefully ruled out. EPS is recommended in symptomatic patients only for the assessment of supraventricular arrhythmia.

2. **Asymptomatic patients** displaying a Brugada ECG (spontaneously or after sodium channel block) should undergo EPS if there is a family history of sudden cardiac death suspected to be due to Brugada syndrome. EPS may be justified when the family history is negative for sudden cardiac death if the Type 1 ST-segment elevation occurs spontaneously. If inducible for ventricular arrhythmia, the patient should receive an ICD. This was recommended as a Class IIa indication for patients presenting with a spontaneous Type I ST-segment elevation and as a Class IIb for patients who display a Type I ST-segment elevation only after sodium block challenge. More recent data (discussed above) have called these recommendations into question and suggest that it might be more appropriate to consider both as Class IIb indications.

Asymptomatic patients who have no family history and who develop a Type 1 ST-segment elevation only after sodium channel blockade should be closely followed-up. As additional data become available, these recommendations will no doubt require further fine-tuning.

Although arrhythmias and sudden cardiac death generally occur during sleep or at rest and have been associated with slow heart rates, a potential therapeutic role for cardiac pacing remains largely unexplored. Haissaguerre and co-workers171 reported that focal radiofrequency ablation aimed at eliminating the ventricular premature beats that trigger VT/VF in the Brugada syndrome may be useful in controlling arrhythmogenesis. However, data relative to a cryosurgical
Table VI.
Indications for ICD Implantation in Patients with the Brugada Syndrome

Spontaneous Type 1 ECG

**Symptomatic**
- Aborted SCD
- Syncope
- Seizure NAR

Evaluate for clear extracardiac cause
- ICD (Class I)
- Close Follow-up

**Asymptomatic**
- Family History of SCD suspected to be due to BS
- No Family History

EPS (Class Ila)
- ICD (Class Ila) Follow-up

EPS justified (Class IIa)
- ICD (Class IIa) Close Follow-up

**Sodium Channel Block-induced Type 1 ECG**

**Symptomatic**
- Aborted SCD
- Syncope
- Seizure NAR

Evaluate for clear extracardiac cause
- ICD (Class I)
- Close Follow-up

**Asymptomatic**
- Family History of SCD suspected to be due to BS
- No Family History

EPS justified (Class IIb)
- ICD (Class IIb) Close Follow-up

EPS recommended for assessment of supraventricular arrhythmias

**Class I:** Clear evidence that procedure or treatment is useful or effective
**Class II:** Conflicting evidence concerning usefulness or efficacy
**Class IIa:** Weight of evidence in favor of usefulness or efficacy
**Class IIb:** Usefulness or efficacy less well established

BS = Brugada syndrome; EPS = electrophysiologic study; NAR = nocturnal agonal respiration; SCD = sudden cardiac death

(from Antzelevitch et al. / with permission)
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Figure 13. Effects of I_{to} blockers 4-AP and quinidine on pinacidil-induced phase 2 reentry and VT in the arterially perfused RV wedge preparation. In both examples, 2.5 mM pinacidil produced heterogeneous loss of AP dome in epicardium, resulting in ST-segment elevation, phase 2 reentry, and VT (left); 4-AP (A) and quinidine (B) restored epicardial AP dome, reduced both transmural and epicardial dispersion of repolarization, normalized the ST segment, and prevented phase 2 reentry and VT in continued presence of pinacidil. From Yan and Antzelevitch with permission.

approach or the use of ablation therapy are very limited at this point in time.

Pharmacologic Approach to Therapy

ICD implantation is the mainstay of therapy for the Brugada syndrome. Although feasible, implantation is challenging in infants and is not an adequate solution for patients residing in regions of the world where an ICD is unaffordable. A pharmacologic solution is desirable as an alternative to device therapy in these cases, as well as in minimizing the firing of the ICD in patients with frequent events.

The quest for a pharmacologic treatment has been focused on rebalancing of the ion channel current active during the early phases of the epicardial AP in the right ventricle, so as to reduce the magnitude of the AP notch and/or restore the AP dome. Table V lists the various pharmacologic agents thus far investigated. Antiarrhythmic agents, such as amiodarone and β blockers, have been shown to be ineffective. Class IC antiarrhythmic drugs (such as flecainide and propafenone) and Class IA agents, such as procainamide, are contraindicated because of their effects to unmask the Brugada syndrome and induce arrhythmogenesis. Disopyramide is a Class IA antiarrhythmic that has been demonstrated to normalize ST-segment elevation in some Brugada patients but to unmask the syndrome in others.

Because the presence of a prominent transient outward current, I_{to}, is fundamental to the mechanism underlying the Brugada syndrome, the most prudent general approach to therapy, regardless of the ionic or genetic basis for the disease, is to partially inhibit I_{to}. Cardioselective and I_{to}-specific blockers are not currently available. 4-aminopyridine is an agent that is ion-channel specific at low concentrations, but is not cardioselective in that it inhibits I_{to} in the nervous system. Although it is effective in suppressing arrhythmogenesis in wedge models of the Brugada syndrome (Fig. 13), it is unlikely to be of clinical benefit because of neurally mediated adverse effects.

An agent currently on the market in the United States and other regions of the world with significant I_{to} blocking properties is quinidine. Accordingly, we suggested several years ago that this agent may be of therapeutic value in the Brugada syndrome. Quinidine has been shown to be effective in restoring the epicardial AP dome, thus normalizing the ST segment and preventing phase 2 reentry and polymorphic VT in experimental models of the Brugada syndrome (Fig. 13). Clinical evidence of the effectiveness of quinidine in normalizing ST-segment elevation in patients with the Brugada syndrome has been reported as well (Fig. 14). Quinidine has also been reported to be effective in suppressing arrhythmogenesis in an infant too young to receive an ICD.

A prospective study of 25 Brugada syndrome patients orally administered quinidine bisulfate (1483 ± 240 mg), reported by Belhassen and Viskin, evaluated the effectiveness of quinidine in preventing inducible and spontaneous VF. There were 15 symptomatic patients (7 cardiac arrest survivors and 7 with unexplained syncope) and 10 asymptomatic patients. All 25 patients had inducible VF at baseline electrophysiological study. Quinidine prevented VF induction in 22 of the 25 patients (88%). After a follow-up period of 6 months to 22.2 years, all patients were alive. Of 19 patients treated with oral quinidine for 6 to 219 months (56 ± 67 months), none developed arrhythmic events. Administration of quinidine
was associated with a 36% incidence of side effects, principally diarrhea, which resolved after drug discontinuation. The authors concluded that quinidine effectively suppresses VF induction as well as spontaneous arrhythmias in patients with Brugada syndrome, and may be useful as an adjunct to ICD therapy or as an alternative to ICD in cases in which an ICD is refused, is unaffordable, or under other circumstances in which ICD implantation is not feasible. These results are consistent with those reported for the same group in prior years and more recently by other investigators. The study by Hermida and co-workers was the first to report the results of prospective, although relatively small, clinical trials. A recent relatively small study by Mizusawa et al. showed that low-dose quinidine (300–600 mg) can prevent electrophysiologic induction of VF and has a potential as an adjunctive therapy for Brugada syndrome in patients with frequent ICD discharges. There is a clear need for a large, randomized, controlled clinical trial to assess the effectiveness of quinidine, preferably in patients with frequent events who have already received an ICD.

There is a clear need for a more cardioselective and Ito-specific blocker as an addition to the limited therapeutic armamentarium currently available to combat this disease. Another agent being considered for this purpose is the drug tedisamil, currently being evaluated for the treatment of atrial fibrillation. Tedisamil may be more potent than quinidine because it lacks the inward current blocking actions of quinidine, while potently blocking Ito. The effect of tedisamil to suppress phase 2 reentry and VT in a wedge model of the Brugada syndrome is illustrated in Figure 15. Quinidine and tedisamil are both capable of suppressing the substrate and trigger for the Brugada syndrome via their inhibition of Ito. Both, however, also block IKr and thus have the potential to induce an acquired form of the long-QT syndrome. Thus, these agents may substitute one form of polymorphic VT for another, particularly under conditions that promote TdP, such as bradycardia and hypokalemia. This effect of quinidine is minimized at high plasma levels, because at these concentrations quinidine block of INa counters the effect of IKr block to increase TDR, the substrate for the development of TdP arrhythmias. High doses of quinidine (1,000–1,500 mg/day) are recommended in order to effect Ito block, without inducing TdP.

Figure 14. Twelve-lead electrocardiogram (ECG) tracings in an asymptomatic 26-year-old man with the Brugada syndrome. (Left) Baseline: Type 2 ECG (not diagnostic) displaying a “saddleback-type” ST-segment elevation is observed in V2. (Center) After intravenous administration of 750 mg procainamide, the Type 2 ECG is converted to the diagnostic Type 1 ECG consisting of a “coved-type” ST-segment elevation. (Right) A few days after oral administration of quinidine bisulfate (1,500 mg/day, serum quinidine level 2.6 mg/L), ST-segment elevation is attenuated in the right precordial leads. VF could be induced during control and procainamide infusion, but not after quinidine. From Belhassen et al. with permission.
Another potential therapeutic candidate is an agent reported to be a relatively selective $I_{to}$ and $I_{Kur}$ blocker, AVE0118. Figure 16 shows the effect of AVE0118 to normalize the ECG and suppress phase 2 reentry in a wedge model of the Brugada syndrome. This drug has the advantage that it does not block $I_{Kr}$, and therefore does not prolong the QT interval or have the potential to...
Figure 17. Effect of dmLSB to suppress the arrhythmogenic substrate of the Brugada syndrome in three experimental models. Phase 2 reentry was induced in three separate models of the Brugada syndrome. Terfenadine (5 μM, A), verapamil (5 μM, B), or pinacidil (6 μM, C) induce heterogeneous loss of the epicardial action potential dome and ST-segment elevation. Phase 2 reentry occurs as the dome is propagated from Epi 1 to Epi 2, triggering either a closely coupled extrasystole or polymorphic ventricular tachycardia. In all 3 models, addition of dmLSB (10 μM) normalizes the ST segment and abolishes phase 2 reentry and resultant arrhythmias. From Fish et al.\textsuperscript{115} with permission.
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References


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192. Myers GB. Other QRS-T patterns that may be mistaken for myocardial infarction. IV. Alterations in blood potassium: myocardial ischemia; subepicardial myocarditis; distortion associated with arrhythmias. Circulation 1956; 2:75–93.


