ABSTRACT: Although there is consensus on the management of patients with Brugada Syndrome with high risk for sudden cardiac arrest, asymptomatic or intermediate-risk patients present clinical management challenges. This document explores the management opinions of experts throughout the world for patients with Brugada Syndrome who do not fit guideline recommendations. Four real-world clinical scenarios were presented with commentary from small expert groups for each case. All authors voted on case-specific questions to evaluate the level of consensus among the entire group in nuanced diagnostic and management decisions relevant to each case. Points of agreement, points of controversy, and gaps in knowledge are highlighted.

**Key Words:** catheter, chest pain, sleep apnea, sudden cardiac arrest, ventricular fibrillation

Brugada syndrome (BrS) is a heritable arrhythmia syndrome associated with ECG features of ST-segment elevation in the right precordial leads followed by T-wave inversion and increased risk of sudden cardiac arrest (SCA) in patients with a structurally normal heart, though microstructural abnormalities are likely present. Patients with BrS may present with syncope or aborted SCA or be asymptomatic. Although there is consensus on the management of patients with BrS with high risk for SCA, asymptomatic or intermediate-risk patients present clinical management challenges. Harnessing experts from around the globe, this document explores the management opinions of clinicians for patients who do not fit guideline recommendations.

Four real-world clinical scenarios, not modified for the purpose of this publication, were presented to small groups of experts, who discuss management recommendations, including points of agreement or disagreement, and compose voting questions for all authors. All authors then voted on case-specific questions to further evaluate the level of consensus among the entire group in nuanced diagnostic and management decisions relevant to each case. Finally, key points as well as gaps in knowledge are summarized (Tables 1 through 3).

**CASE 1**
A 50-year-old man with bipolar disorder (stable off medication), hypertension, and obstructive sleep apnea presented to the emergency department with myalgias, nausea, headache, fatigue, and chest pain. He was febrile at 38.6°C. Serum electrolytes and cardiac enzymes were normal. ECG showed ST elevation in V1 to V3 (Figure 1A). Urgent coronary angiography showed no significant obstructive coronary artery disease. However, during the procedure, the patient had ventricular fibrillation...
fibrillation (VF) requiring defibrillation. This VF did not occur during contrast injection or catheter manipulation with the catheter resting in the aorta.

Repeat ECG when the patient was afebrile showed resolution of ST elevation in V1 to V3 (Figure 1B). Specific and expanded genetic testing was negative for pathogenic variants. Cardiac imaging (echocardiogram and magnetic resonance imaging) was normal. Ventricular programmed electrical stimulation using up to 3 extra-stimuli repeatedly induced self-terminating ventricular tachycardia (15–20 seconds).

KEY QUESTIONS

(1) Would you perform a drug provocation study or do additional risk stratification? (2) Would you recommend an implantable cardioverter defibrillator (ICD)? (3) What type of screening/counseling would you recommend to family members given negative genetic testing?

EXPERT PANEL COMMENTARY (CERRONE [CHAIR], WILDE, LONDON, BEHR, SHIMIZU)

The majority of panelists agreed that based on the data provided, including fever-induced type 1 ECG pattern and nonprovoked VF, the diagnosis of BrS is confirmed. However, one panelist suggested the diagnosis could be questionable based on the Shanghai score system, in which this individual would reach only 3 points, 3.5 points considered diagnostic, since a type 1 ECG during fever is not classified as spontaneous. The possibility of a false positive was also raised, based on the morphology of the ST segment elevation in V1.

All panelists agreed on the known limitations of a drug challenge and the possibility of false positive results. A pharmacological challenge was considered unnecessary for the diagnosis since the patient already showed a fever-induced type 1 pattern, although some experts suggested that a provocative test could validate the one-time finding of type 1 during fever and be used as a tool for cascade screening.

Most of the expert panel for case 1 supported additional screening tools including a high-lead ECG (V1 and V2 placed in the second and third intercostal space) and 12-lead Holter (with the option of recording high and conventional precordial leads simultaneously). Additional components to help risk stratification and overall assessment include detailed medical and family history, reviewing past ECGs, signal-averaged ECG, treadmill exercise testing, and careful analysis of ECG characteristics such as QRS spike wave at V1 to V3 leads, J wave at inferolateral, QRS duration, and 12-lead Holter.

The exact role of provocative drug testing in relatives of patients with BrS phenocopy. Value of genetic testing in the absence of a confirmed Brugada Syndrome diagnosis but with syncope and whether such testing, if performed, should be narrow (SCN5a only) or broad.

Table 2. Points of Controversy

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<td>Electrophysiology study to guide risk stratification in patient with Brugada Syndrome.</td>
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Table 3. Gaps in Knowledge

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<td>The exact role of provocative drug testing in relatives of patients with BrS (when and how often).</td>
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<td>The role of asymptomatic fever-induced type 1 ECG pattern without spontaneous type 1 ECG pattern, family history of BrS, or a P/LP mutation in a BrS susceptibility gene is unresolved.</td>
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<td>The role of provocative sodium channel blocker drug testing to diagnose BrS syndrome. Specifically, additional research is needed to determine the indications and to better define differences in the sensitivity and specificity of different sodium channel blockers and the impact of regional differences in access to certain IV sodium channel blocker drugs.</td>
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BrS indicates Brugada Syndrome; ICD, implantable cardioverter defibrillator; and VF, ventricular fibrillation.
The critical question to consider when deciding on whether to implant an ICD is whether the patient's cardiac arrest was triggered by the angiogram or purely coincidental. If the arrest is considered a spontaneous event, the long-term risk of SCA would be sufficiently high (around 8%/y) to support consideration of an ICD. In addition, none of the currently available risk stratification strategies have sufficient negative predictive value given a history of unprovoked SCA. The experts who leaned toward defining the VF as a nonspontaneous episode, because it occurred during a medical procedure (even if not connected to high-risk interventions), were inclined to not implant an ICD. The experts emphasized the need for a detailed discussion with the patient regarding risks and benefits and gaps of knowledge. The possibility of long-term monitoring with an ILR should be considered if an ICD is not implanted.

If in the future, the patient requires a therapy for Bipolar disease, with drugs that are contraindicated in the setting of BrS (www.brugadadrugs.org), all authors agree

Figure 1. Case 1 ECGs.
A, Spontaneous ECG during fever. B, Repeat ECG when afebrile.
that he should be followed closely with ECG monitoring at regular intervals. There was agreement on screening relatives with ECG, high-lead ECG, and 12-lead Holter looking for spontaneous type 1 pattern.\(^{17}\) Relatives should be counseled to have an ECG recorded during a fever when possible and to implement modifying measures, such as prompt treatment of fevers and avoidance of agents known to be proarrhythmic in the setting of BrS.\(^{2,3,13}\) Children should repeat an ECG after puberty, and all adult family members with a first negative ECG should repeat it in several years. Provocative drug challenge was not deemed necessary but could be considered, after informed decision with the relatives, especially in the presence of a suspicious ECG pattern.

**CASE 1 SUMMARY (PEREZ, RODEN)**

The experts agreed on most of the major management questions raised by this case of a 50-year-old man with a type I Brugada pattern in the setting of a febrile illness and a cardiac arrest during coronary angiography (Figure 2; Table S1). However, there are a few points of disagreement worth highlighting. While the 2013 expert consensus statement on inherited arrhythmias states that BrS “is diagnosed in patients with ST-segment elevation with Type 1 morphology…either spontaneously or after provocative drug test…,”\(^{18}\) a minority of the experts here proposed that a type 1 pattern induced by a fever should not be considered spontaneous and that there is a role for drug provocation in this case. However, there is a gap in knowledge of whether a sodium channel blocker test is more accurate than a fever-induced ECG in attesting a BrS diagnosis. Further research is needed to answer this question. The disagreement on appropriateness of EPS and ICD was based in part on whether the episode of VF was provoked by the coronary angiogram of which most of the expert commentary panel for Case 1 felt it was unprovoked. The entire author panel was more likely to recommend an ICD. Regardless, there was agreement that if the VF episode was unprovoked, then the EPS would not add value as a negative study would not be sufficiently reassuring. Finally, there was discrepancy on the ideal strategy for family screening, possibly due to the lack of a clear guideline on routine use of provocative drug testing to screen relatives and the frequency of screening. While most experts agreed a drug challenge should be considered in family members if there is clinical suspicion, it remains controversial whether routine drug challenge should be recommended in the absence of symptoms or equivocal ECG findings.

**CASE 2**

A 33-year-old female was seen in the emergency department following her first episode of syncope preceded by a prodrome of lightheadedness (ECG; Figure 3A). She reported symptoms of an upper respiratory infection but no fever. She had no significant past medical history and her family history was positive for coronary artery disease. Genetic testing revealed a pathogenic mutation in SCN5A (c.2533delG). At the time of expert consultation, the ECG (Figure 3A) from the emergency department

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**Figure 2. Case 1 group voting.**

Survey results from questions related to case 1. ICD indicates implantable cardioverter defibrillator.
was not immediately available and an in-office ECG (Figure 3B) was obtained without a type 1 Brugada ECG pattern. As such, she underwent procainamide challenge and developed type 1 Brugada ECG pattern.

**KEY QUESTIONS**

(1) Would you perform a diagnostic EPS? (2) Would you recommend an ICD? (3) Should asymptomatic relatives with normal clinical evaluation and negative genetic testing be released from clinical follow-up?

**EXPERT PANEL COMMENTARY (SY [CHAIR], DEASMUNDIS, GOLLOB, KRAHN, SARQUELLA-BRUGADA)**

Case 2 expert panelists agreed that the initial ECG from the emergency department shows a spontaneous type 1 BrS ECG pattern. Procainamide challenge reproduced the ECG pattern, but the provocation study was considered unnecessary given the index ECG. The presence of a pathogenic variant in SCN5A further confirms the diagnosis and may have implications in terms of arrhythmic
outcomes. The panel agreed that the interpretation of genetic variants is ideally performed in a multidisciplinary cardio genetics service, especially in light of recent evidence that non-SCN5A variants often have limited/disputed evidence for pathogenicity in BrS.

In terms of diagnosis, the patient has BrS based on her ECG, clinical presentation, and genetic testing result. This yields a Shanghai score of at least 5.0, confirming BrS diagnosis.

Although arrhythmic events dominate in males >40 with BrS, this younger female patient's history of syncope is concerning and her prognosis and management hinges on the evaluation of the syncopal event. Systematic history-taking is crucial in differentiating nonarrhythmic syncope and arrhythmic syncope. Specifically, the presence of prodromal symptoms (nausea, diaphoresis, cough, or micturition) would point toward nonarrhythmic cause. Importantly, nonarrhythmic syncope occurs frequently in patients with BrS (up to 57% in 1 study) but is not associated with malignant outcomes. In contrast, patients with arrhythmic syncope have a risk of subsequent SCA.

Clinical history should provide sufficient discrimination of the likely mechanism of syncope, and additional investigations such as tilt table testing and EPS are nonspecific.

Risk stratification in BrS continues to evolve. The presence of a spontaneous type 1 Brugada ECG pattern is a consistent marker of increased risk, especially in the setting of syncope. Beyond a spontaneous type 1 Brugada ECG pattern, additional ECG markers have been reported to be associated with an increased risk of arrhythmia but these were not present in this patient.

The utility of an EPS for risk stratification remains contentious. In particular, the incremental value of VF inducibility is questionable in the present case if the patient is deemed to have arrhythmic syncope because a negative test would be insufficient to withhold recommending an ICD. Hence, 3 panelists would not recommend an EPS in this scenario. However, 2 panelists would recommend an EPS to further evaluate the arrhythmic risk and evaluate the HV interval and sinus node recovery time given that brady arrhythmias may be associated with BrS, especially in patients with a pathogenic SCN5A variant.

There is complex interplay between gender and risk in BrS. Male patients with BrS are more likely to exhibit a spontaneous type 1 BrS pattern as well as inducibility of VF and have a greater risk of malignant arrhythmia. The present patient poses a less common clinical scenario, a female patient with a spontaneous type 1 BrS pattern and a pathogenic mutation in SCN5A. Epidemiological data suggest that gender alone is not an independent predictor of outcomes once other variables such as the presence of a spontaneous type 1 ECG pattern are taken into account. Moreover, female BrS patient with pathogenic SCN5A mutations may have an increased risk of malignant arrhythmias. However, it is acknowledged that risk stratification in female patients with BrS is less well understood because the vast majority of patients in clinical studies, and even more so in those with clinical events, are male.

Current guidelines would recommend the consideration of a prophylactic ICD in the setting of probable arrhythmic syncope and a spontaneous Type 1 ECG pattern. However, it is critical to engage the patient in shared decision-making after a thorough discussion of the potential benefits as well as the lifetime risks of ICD implantation in young patients including infection, system revision, and inappropriate shocks. If the patient declines ICD implantation, the merits and limitations of adjuvant strategies such as quinidine therapy and catheter ablation can be discussed as alternatives with limited evidence from small observational studies.

Lifestyle advice regarding medication avoidance (brugadadrugs.org), restraint from alcohol intoxication, and prompt fever treatment is recommended.

The experts agreed that the patient's relatives should be offered clinical evaluation as well as cascade testing for the pathogenic SCN5A variant identified in the proband. Relatives who have clinical evidence of BrS and carriers of the SCN5A variant should be carefully screened for arrhythmic symptoms. Sodium-channel blocker challenge can be offered in selected patients based on their symptom status, ECG, and preference. Asymptomatic patients should receive lifestyle advice and clinical follow-up. A diagnostic EPS is not recommended in asymptomatic relatives.

In general, asymptomatic relatives with a completely normal resting ECG (including high-lead ECG) and negative for the SCN5A variant can be released from clinical follow-up. However, there is an increasing appreciation of the complex heritability of BrS. Of note, a patient's genetic background (beyond SCN5A variants) may contribute to variable expressivity in families with a pathogenic variant in SCN5A.

CASE 2 SUMMARY (CUTLER, HUANG)

This case identifies key questions: is provocative drug testing needed in a patient who presents with a spontaneous type 1 pattern ECG? The panelists agreed that the presenting ECG displayed a Type 1 pattern and that a procainamide challenge test was not needed. Was the syncopal episode arrhythmic or nonarrhythmic? The panel was unanimous in recommending a detailed history of the syncope to distinguish arrhythmic versus nonarrhythmic syncope as the determination of arrhythmic syncope is crucial in the decision to recommend an ICD. Finally, does an EPS add value to the risk stratification of this patient? As in the literature, whether an EPS should be performed was debated. Three of the 5 panelists...
would not recommend EPS implying that syncope with ECG findings was sufficient for diagnosis and prescribing treatment. In contrast, 2 panelists recommended EPS looking for ventricular arrhythmia inducibility and SA nodal or AV conduction pathology.

The entire author panel was divided on whether to recommend an EPS; the majority would not (Figure 4; Table S2). While 74% of the experts would recommend an ICD implant, an additional 18.5% would recommend an ICD if an EPS was positive. There was consensus that asymptomatic relatives with negative genetic testing could be released from follow-up.

In conclusion, case 2 highlights the importance of a thorough history to distinguish between arrhythmic and nonarrhythmic syncope. However, there are instances when all available clues still may not clearly differentiate arrhythmic versus nonarrhythmic syncope and additional risk stratification tools, for example, spontaneous versus induced ECG changes, genotype details if positive, or EPS may be needed.

CASE 3

A 26-year-old female became unresponsive following a period of diaphoresis, flushing, and tunnel vision after ingesting alcohol. A bystander applied an automated electrical defibrillator and no shock was advised. When emergency medical services arrived, the patient was arousable to sternal rub. In the emergency room, an ECG (Figure 5A) and cardiac imaging (echocardiogram, cardiac magnetic resonance imaging) were normal. Family and personal history were negative for SCA, syncope, or febrile seizures.

Subsequently, a procainamide challenge was performed (Figure 5B and 5C). She was discharged home with a life vest and had syncope while wearing the life vest. No ventricular arrhythmia was detected on life-vest interrogation. Genetic testing showed a likely pathogenic mutation in SCN5A (c.4978A>G). A diagnostic EPS was performed with no inducible ventricular arrhythmias.

**KEY QUESTIONS**

1. Would you diagnosis this case as BrS? (2) Would you perform additional risk stratification and recommend an ICD? (3) Would you recommend ECG screening and genetic testing for family members?

**EXPERT PANEL COMMENTARY (CROTTI [CHAIR], ARBELO, BRUGADA, SACHER, WATANABE)**

All experts agreed that the ECGs provided do not fulfill criteria for the diagnosis of BrS. However, they acknowledged that the diagnosis cannot be ruled out completely for the following reasons: (1) the absence of a high-lead ECG at baseline and during procainamide challenge and (2) use of procainamide as a drug challenge instead of ajmaline.

BrS can only be diagnosed in the presence of the type 1 Brugada pattern characterized by J point elevation of >2 mV with coved ST elevation and T wave inversion in at least 1 right precordial ECG lead (ie, V1 or V2). Placement of the right precordial leads in a more superior position (ie, second or third intercostal spaces) increases the ECG sensitivity to identify a type 1 Brugada ECG pattern. Other situations such as fever, vagal stimulation, alcohol, cocaine intoxication, or electrolyte abnormalities may unmask type 1 pattern when ECG manifestations are not apparent at baseline. The presence of other known causes of ST-segment elevation in right precordial leads (so-called phenocopies) should be excluded.

When the baseline ECG does not show a typical type 1 Brugada ECG pattern, but there is a reasonable suspicion,
intravenous administration of a sodium channel blockers may convert the ECG pattern into Type 1.53–55 Unfortunately, not all drugs seem to have the same diagnostic yield for drug-induced Type 1 ECG pattern. In retrospective analysis, Ajmaline may be superior to other sodium channel blockers; yet, the sensitivity and specificity of provocative drug testing remain elusive and Ajmaline is not available in all countries.56–58 As such, further research is needed to better define the role of provocative drug testing with IV sodium channel blockade in the diagnosis of BrS.

The appropriate strategy for risk stratification depends on whether the diagnosis of BrS is confirmed. If a diagnosis of BrS is not made, the workup and management should follow the recommendations for the management of patients with syncope.59 If BrS is confirmed, a detailed evaluation of each syncopal event is warranted60 in an attempt to classify each as arrhythmic or not25,61 as recommended arrhythmic syncope treatment in BrS is an ICD.13,18,35,62 This is particularly important given the high prevalence of vasovagal syncope in published cohorts of BrS.61 Based on the clinical information available for this case, including the absence of ventricular arrhythmia on AED and LifeVest, this patient’s syncope was likely not arrhythmic.

In patients with BrS with syncope of unclear mechanism an EPS to assess inducibility of sustained (ie, ≥30 seconds) polymorphic ventricular arrhythmias could be appropriate because the EPS has also shown to have a negative predictive value (92.4%).61 However, in this case we do not have a diagnosis of BrS and therefore, the majority of panelists would not recommend an EPS. Three experts think that in the absence of a BrS diagnosis, without personal history of febrile seizures or palpitation, no family history of SCA, and nonarrhythmic syncope, only regular clinical follow-up is recommended.18,35,62 Two experts would recommend an implantable loop recorder, and all panelists agreed that an ICD was not indicated.

According to a recent consensus document on the use of molecular screening in cardiac diseases, genetic testing should be performed only in patients with a type 1 standard or high-lead ECG occurring either spontaneously or induced by sodium-channel block, and only SCN5A should be screened in a clinical setting since it is the only gene with definite association with BrS.38,63 All panelists agreed that in the absence of a diagnosis of BrS, molecular screening should not have been performed. However, once a likely pathogenic variant on SCN5A was identified, the data should be appropriately managed.

Two experts suggested that cascade screening should not be performed unless a type 1 pattern was identified in the proband. In contrast, 2 panelists recommended that
genetic and complete clinical evaluation should be offered to first-degree relatives. The remaining expert recommended that variant classification should be reevaluated in an independent laboratory with a specific expertise. Importantly, the distinction between variant of uncertain significance and likely pathogenic can sometimes be subtle and change over time. Indeed, this variant has been reclassified as a variant of uncertain significance (PP3-PP5-BS2) in an independent laboratory and should not be used to support diagnosis or for cascade screening.

**CASE 3 SUMMARY (PROBST, LUBITZ)**

In the present case, a young adult woman experienced syncope after a brief prodrome. An AED was applied, and no shock was advised. The patient was rousable without a shock, suggesting a nonarrhythmic event. Her subsequent syncopal event while wearing a LifeVest confirmed her lack of tachyarrhythmia. Her ECG showed transient abnormal early repolarization in the right precordial leads, reproduced with a procainamide challenge.

Of the entire author voting group, most (70%) indicated that they would not have pursued EPS. This observation highlights the variability of opinion and practice among providers of the utility of an electrophysiology study in the diagnostic workup of unexplained syncope, particularly in the setting of confounding genetic testing results, although such testing may not have been indicated. Nevertheless, EPS was performed and was negative (Figure 6; Table S3). At variance to current consensus documents, slightly over half the experts would have performed genetic testing, with 41% favoring a broad genetic panel and 11% focusing on SCN5A variants only. Genetic testing identified a likely pathogenic missense variant in SCN5A (c.4978A>G), later reclassified as a variant of uncertain significance. Most (63%) of experts were in favor of the implantation of a loop recorder to further assess the syncope cause.

The current case highlights the importance of understanding the cause of syncope, diagnostic electrocardiographic criteria for BrS, and genetic variant interpretation to avoid unnecessary exams and potentially harmful treatments for patients.

**CASE 4**

A 32-year-old man without known past medical history was arrested for operating a vehicle while intoxicated. He was found with a bag containing a white substance in his mouth that apparently burst. While in police custody, he had a seizure treated with midazolam and then had a VF cardiac arrest. He was hypotensive and required intubation by the emergency medical service. In the emergency department, telemetry monitoring showed marked ST elevation. ECG was performed (Figure 7A) and urgent EP consult requested for rule out BrS. Troponin was elevated, blood alcohol was 56 mg/dL, and the toxicology screen was positive for cocaine. Within 3 hours of presentation to the hospital, his ECG (Figure 7B) normalized and respiratory status improved. He was extubated, became agitated, and left the hospital against medical advice.

**KEY QUESTIONS**

1. If he had recurrent episodes of VF in the emergency department, how would you have managed this? (2) If his family brought him to your clinic for a follow-up visit, what further testing, if any, would you recommend? (3) Can cocaine and alcohol intoxication be considered like performing an ajmaline or procainamide challenge with respect to diagnosis of BrS?
EXPERT PANEL COMMENTARY (EXPERTS: MACKALL [CHAIR], NADEMANEE, SCHEINMAN, SHOEMAKER)

The patient presented with VF arrest in the setting of cocaine and alcohol intoxication. In addition to VF, acute cocaine intoxication can present with acute hypertension, coronary vasospasm, and myocardial infarction/ischemia. Moreover, chronic cocaine abuse increases risk of acute coronary syndrome, cardiomyopathy, and may increase risk of coronary artery disease. The presenting ECG shows prolonged PR interval and QRS complex and ST segment elevation with T wave inversion consistent with possible Brugada pattern or cardiac ischemia. One expert felt the ECG more likely reflected a conduction...
block due to sodium channel intoxication than a Type 1 Brugada pattern.65

Cocaine intoxication is a recognized clinical scenario in which the Brugada pattern ECG represents a Brugada phenocopy. Brugada phenocopies have been described in other cases of overdose with medications that have sodium channel–blocking effects such as tricyclic antidepressants, anti-seizure drugs, or Class IC antiarrhythmics.66 The electrophysiological effects of cocaine are related to sodium channel blockade, manifesting as prolonged PR interval and QRS widening. This patient had both cocaine and alcohol intoxication which is more toxic than cocaine alone because the metabolite cocaethylene has a more pronounced sodium channel effect and longer half-life.67 The primary difference between drug-induced Brugada ECG and Brugada phenocopy is the presumed level of sodium channel blockade with a therapeutic dose of a sodium channel blocker compared with drug overdose.

The treatment of recurrent VF in this patient should include prompt defibrillation followed by evaluation and treatment for acute coronary vasospasm or myocardial infarction, as appropriate. Fluid resuscitation and sodium bicarbonate are recommended to treat acidosis and restore sodium channel function by promoting dissociation of cocaine from the sodium channels. Furthermore, cocaine toxicity can result in QT interval prolongation secondary to blocking of potassium channels, leading to Torsade de Pointes. In such cases, the preferred treatment would include IV magnesium and lidocaine. Importantly, isoproterenol and beta-blockers are contraindicated with recurrent VF from cocaine toxicity.

Because the ECG was not diagnostic for BrS, panelists disagreed on whether isoproterenol would be the medication of choice. Quinidine would not be recommended for treating recurrent VF because of its sodium channel–blocking properties and risk for hypotension. One of the key variables missing from the patient's summary is the body temperature on arrival at the emergency department. Cocaine toxicity causes hyperthermia, which greatly affects Brugada substrates and could precipitate tachyarrhythmia.

Brugada phenocopies have been described in clinical settings other than drug overdose, including electrolyte disorders and inflammatory syndromes. Distinguishing phenocopy from BrS involves taking a careful family history of SCA and a personal history of syncope or febrile seizure. A review of pertinent laboratory data to evaluate for electrolyte disturbances and current medications to identify drugs that potentiate sodium channel blockade (eg, lithium or tricyclic antidepressants, and phenytoin) should be performed. On physical examination, the presence of pectus excavatum68 or pericardial rub (pericarditis)69 should be noted, if present, as both conditions may present with a type 2 Brugada ECG pattern. A high-lead ECG should be performed, and prior ECGs should be reviewed to verify the absence or presence of spontaneous type 1 Brugada pattern. Additional imaging is suggested to rule out any structural cardiac condition or pulmonary embolism.70 One expert observed that these data would be helpful in a decision regarding genetic testing.

Two panelists asserted that if the office evaluation was negative, then Brugada phenocopy in the setting of cocaine intoxication was likely. Two experts would perform a sodium channel blocker challenge as a negative drug challenge would confirm the diagnosis of Brugada phenocopy.51,71 Genetic testing would only be considered if a BrS diagnosis was possible or probable based on the Shanghai Score, acknowledging that SCN5A mutations are identified in only 20% of cases.22,73

Cocaine with alcohol intoxication cannot be considered the equivalent of an ajmaline or procainamide challenge. The levels of cocaine and alcohol and their metabolite coca-ethylene contribute to metabolic derangement, altered sympathetic and parasympathetic activity, and sodium channel blockade. ECG changes demonstrating a wide QRS complex, a Brugada pattern and ventricular arrhythmias have all been reported in cocaine intoxication due to primarily sodium channel blocking effects. In contrast, a positive drug challenge with ajmaline or procainamide results in a type I pattern that reflects abnormal sodium channel function at doses that would not normally elicit ECG changes. While the experts agreed that a Brugada pattern evident with cocaine intoxication would not be diagnostic of BrS, 1 panelist felt that a Brugada pattern in the presence of alcohol intoxication with an otherwise negative toxicology screen would be equivalent to a drug challenge.

**CASE 4 SUMMARY (HORIE, KAUFMAN)**

Case 4 is challenging because this young patient had a cardiac arrest in the setting of cocaine and alcohol intoxication, then left the hospital against medical advice and was not available for further evaluation. The experts agreed on the details of acute management. If the patient were available for further evaluation, the experts would focus on personal and family history, examination for pectus excavatum or pericardial rub, and additional ECG recordings. They would also consider follow-up imaging (echocardiogram or magnetic resonance imaging), and drug challenge to distinguish BrS phenocopy from actual BrS. Genetic testing would be considered only if diagnosis of probable BrS was made. The authors agreed that BrS phenocopy induced by cocaine was not equivalent to ajmaline or procainamide drug challenge. One expert considered that a BrS pattern induced by alcohol alone would be equivalent to a drug challenge.

When the entire group of authors was polled, there were different opinions on whether to consider an ICD (Figure 8; Table S4). The majority (74%) said no, while the others would implant an ICD either based on the VF arrest alone (there was concern for likelihood of repeat
drug exposure) or if EPS or imaging studies were abnormal. The authors were divided on whether to proceed with drug challenge to diagnose BrS and not simply BrS phenocopy, with 63% in favor. Most authors recommended imaging (echo or magnetic resonance imaging) to identify possible occult structural heart disease that can underlie cardiac arrest in a young person, even if a provocative event is the trigger. One important gap in knowledge is the natural history of cardiac arrest attributed to BrS phenocopy in the setting of substance abuse, and whether the risk of recurrent VF justifies ICD implantation.

CONCLUSIONS
Experts agree that the diagnosis of BrS requires careful evaluation of available clinical history and data to rule out Brugada phenocopy and confirm BrS. Examination of all available ECGs, including high-lead ECGs, is valuable. Once a diagnosis of BrS is confirmed risk stratification is paramount to guide when lifestyle modification is insufficient and ICD implantation, with its serious implications, should be recommended. To this end, it is crucial for the clinician to distinguish arrhythmic from nonarrhythmic syncope. The experts are divided on the best use of additional risk stratification strategies. Focused genetic testing can be appropriate for diagnosed BrS patients and facilitate cascade family screening but is best performed in a multidisciplinary cardigenetics center. Application of the guidelines to real patients requires a thoughtful and individualized approach.

ARTICLE INFORMATION
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REFERENCES

Supplemental Material
Tables S1–S4

Sources of Funding
Dr Arbelo receives support from Fundación La Marato de TV3 (Project 245/JU/2020). Dr Behr receives support from The Robert Lancaster Memorial Fund. Dr Krahn receives support from the Paul Brunes Chair in Heart Rhythm Disorders (Vancouver, BC) Dr London receives support from National Institutes of Health (NIIH) R01 HL065300, R01 HL077396, R01 HL119955. Dr Lubit previously received support from NIH grants R01 HL139731 and R01HL157635, and American Heart Association 18SFRN34250007 during this project. Dr Roden receives support from NIH grants R01 HL149926 and R01HL164675. Dr Eckhardt receives support from NIH grants R01HL163987, R01HL143143, R01HL139738, and the Gary and Mary Weiner Professor of Cardiovascular Medicine Research.

Disclosures
Dr Lubit is a full-time employee of Novartis Institute of BioMedical Research as of July 18, 2022. Dr Lubit previously received sponsored research support from Bristol Myers Squibb, Pfizer, Boehringer Ingelheim, Fitbit, Medtronic, Premier, and IBM and has consulted for Bristol Myers Squibb, Pfizer, Blackstone Life Sciences, and Invitae.

Acknowledgments
Dr Arbelo, Crotti, DeAsmundis, Probst, Sacher, Sanquella-Brugada, and Wilde are members of the European reference Network for rare, low prevalence and complex diseases of the ERN-GUARD Heart.

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Circ Arrhythm Electrophysiol. 2024;17:e012072. DOI: Circulation: Arrhythmia and Electrophysiology January 2024 13

Preparation: American Heart Association


