BRUGADA SYNDROME

EXPERT COMMENTARY

Evaluating the Impact of Sex and Gender in Brugada Syndrome

KEYWORDS. Brugada syndrome, gender, sex, testosterone, transgender.

Drs. Wilde and Postema discuss

The Brugada syndrome (BrS) case described by Drs. Sichrovsky and Mittal is a unique case that may help us to understand the reason for the male predominance of BrS. Before going into the details, it is important to emphasize that this case is another example of the difficulties that apparently still exist in recognizing a BrS electrocardiogram (ECG) pattern. An ECG like this one should not be missed anymore!

The patient described is a genetically female patient who had been using testosterone for about 20 years “to live the life of a transgender male.” He presented with an out-of-hospital cardiac arrest and a BrS type 1 pattern ECG (note, the ECG in Sichrovsky et al.’s Figure 2 is from five months before the arrest). Unfortunately, we do not have an ECG from before the start of the intramuscular testosterone injections. Importantly, the claim that testosterone converted this female into a symptomatic BrS male, the essence of this case report and the basis for the nice subtitle “Gender trumps sex as a risk factor,” can of course only with confidence be made with the demonstration of a normal ECG prior to the testosterone therapy. Yet, the odds are in favor of the interpretation presented by the authors. Indeed, testosterone serum levels have been shown to impact the degree of right precordial ST-segment elevation. At this point, we can say that all interventions that increase the early potassium currents, and testosterone may be one of them, also impact in a negative way the safety of conduction. Hence, the effects of testosterone may also be explained by further deterioration of conduction in the right ventricular outflow tract (RVOT) area.

The authors clearly adhere to the “repolarization theory” as the pathophysiological mechanism of the right precordial ST-segment elevation. At this point, we can say that all interventions that increase the early potassium currents, and testosterone may be one of them, also impact in a negative way the safety of conduction. Hence, the effects of testosterone may also be explained by further deterioration of conduction in the right ventricular outflow tract (RVOT) area.

Finally, we assume that the ectopy shown in this patient is not related to the BrS substrate. Although the origin is in the RVOT area, the coupling interval of ventricular fibrillation (VF)–triggering episodes in BrS patients is shorter as compared with the ectopy in this patient. Earlier studies report a coupling interval of less than 400 ms and, here, it is 440 ms to 560 ms (Figure 2 by Sichrovsky et al.). Also, the fact that quinidine was not effective in suppressing the ectopy (while it is very effective in suppressing more serious arrhythmias, as has been described previously in BrS) is in favor of there being a different mechanism for the patient’s ectopy. This potentially explains why the ablation procedure from the endocardial side was successful, whereas the substrate for BrS-related arrhythmias is expected in the epicardial layer. It is also possible that ablation from the endocardial side does affect the epicardial layer of the RVOT, which, after all, is relatively thin.

In summary, the use of testosterone in this patient most likely contributed to the BrS phenotype and underscores the fact that gender indeed impact the phenotype. The RVOT ectopy is presumably unrelated but may serve as a trigger in the setting of a vulnerable substrate in the epicardial layer of the RVOT region.
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References

1. Sichrovsky TC, Mittal S. Brugada syndrome unmasked by use of testosterone in a transgender male: gender trumps sex as a risk factor. J Innov Cardiac Rhythm Manage. 2019;10(2):3526–3529.
2. Gottschalk BH, Anselm DD, Brugada J, et al. Expert cardiologists cannot distinguish between Brugada phenocopy and Brugada syndrome electrocardiogram patterns. Europace. 2016;18(7):1095–1100.
3. Matsuo K, Akahoshi M, Seto S, Yano K. Disappearance of the Brugada-type electrocardiogram after surgical castration: a role for testosterone and an explanation for the male preponderance. Pacing Clin Electrophysiol. 2003;26(7 Pt 1):1551–1553.
4. Ezaki K, Nakagawa M, Taniguchi Y, et al. Gender differences in the ST segment: effect of androgen-deprivation therapy and possible role of testosterone. Circulation J. 2010;74(11):2448–2454.
5. Shimizu W, Matsuo K, Kokubo Y, et al. Sex hormone and gender difference—role of testosterone on male predominance in Brugada syndrome. J Cardiovasc Electrophysiol. 2007;18:415–421.
6. Wilde AAM, Postema PG, Diego JM, et al. The pathological mechanism underlying Brugada syndrome: depolarization versus repolarization. J Mol Cell Cardiol. 2010;49(4):543–553.
7. Kakishita M, Kurita T, Matsuo K, et al. Mode of onset of ventricular fibrillation in patients with Brugada syndrome detected by implantable cardioverter defibrillator therapy. J Am Coll Cardiol. 2000;36(5):1646–1653.
8. Viskin S, Wilde AA, Tan HL, Antzelevitch C, Shimizu W, Belhassen B. Empiric quinidine therapy for asymptomatic Brugada syndrome: time for a prospective registry. Heart Rhythm. 2009;6(3):401–404.
9. Nademanee K, Veerakul G, Chandanamattha P, et al. Prevention of ventricular fibrillation episodes in Brugada syndrome by catheter ablation over the anterior right ventricular outflow tract epicardium. Circulation. 2011;123(12):1270–1279.

Dr. Brugada remarks

To honor the magnificent contributions of the late Philippe Coumel in the understanding of the mechanisms of cardiac arrhythmias, I coined the term “Coumel’s triangle.” Coumel’s philosophy on cardiac arrhythmias was based on the concept that a complex situation (a cardiac arrhythmia) is likely the result of multiple causation: the interaction between the substrate of the arrhythmia with the modulating factors facilitates the start of the arrhythmia by way of the appropriate triggers. A presented example can be used to clarify these mechanisms, as follows: a patient with Wolff–Parkinson–White (WPW) syndrome is born with an accessory atrioventricular (AV) pathway (the substrate). The arrhythmia, circusmovement tachycardia, can be triggered by extrasystoles or junctional rhythm during bradycardia. However, the arrhythmia will only initiate and perpetuate if the conduction properties of the different pathways involved in reentry are appropriate. The appropriateness of the conduction properties is modulated, among other ways, by the autonomic nervous system. An application of the Coumel’s triangle in the case of WPW syndrome is shown in Figure 1.

Coumel’s triangle can be used to understand cardiac arrhythmias and, in fact, it can be applied to understand any phenomenon or event in any domain of knowledge. Figure 2 shows another succinct application of Coumel’s triangle to understand a gas explosion in a room.

In the present issue, Drs. Sichrovsky and Mittal present a phenomenal case of a female-born patient with BrS who has been living as a transgender male via the chronic use of testosterone. According to their description, the patient developed a BrS ECG pattern under this medication. Subsequently, his BrS pattern turned into a full-blown symptomatic syndrome with repeated cardiac arrest. Appropriate protection with an implantable cardioverter-defibrillator was provided to the patient, together with variable quinidine therapy. Symptomatic frequent ventricular extrasystoles prompted endocardial ablation at the RVOT, with a subsequent event-free follow-up. Based on these observations, the authors concluded that “gender trumps sex when it comes to arrhythmia risk in … Brugada syndrome.” Looking at this case with the glasses of Coumel’s triangle, however, I think it is important to state that I disagree with this statement. Biological sex (sex—the substrate) and
social sex (gender—a modulating factor) do not compete against each other in importance to trigger VF in BrS. Rather, both are part of the same triangle and are equally relevant (Figure 3). In terms of the triggers, I do believe that the extrasystoles (and the bradycardia) were examples in this patient. Elimination of the ventricular extrasystoles with ablation probably eliminated one of the triggers; however, it is also possible that the authors inadvertently also eliminated the BrS substrate. Successful endocardial (instead of epicardial) ablation of BrS substrate has been reported.3

This observation is not surprising, given the very small thickness of the RVOT. Endocardial or epicardial ablation in that area most likely always results in a transmural lesion.3 Drs. Sichrovsky and Mittal should be congratulated by their extraordinary observation. This case makes it also clear that testosterone has to be added to the list of drugs to be avoided in confirmed or suspected BrS (www.brugadadrugs.org).

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References
1. Brugada P, Wellens H. Abstracts: 20 years of programmed electrical stimulation of the heart. June 1-5, 1987, Maastricht, The Netherlands. Pacing Clin Electrophysiol. 1987;10(3 Pt 1): 591–622.
2. Sichrovsky TC, Mittal S. Brugada syndrome unmasked by use of testosterone in a transgender male: gender trumps sex as a risk factor. J Innov Cardiac Rhythm Manage. 2019;10(2):3526–3529.
3. Tauber PE, Mansilla V, Brugada P, et al. Endocardial approach for substrate ablation in Brugada syndrome: epicardial, endocardial or transmural substrate?. J Clin Exp Res Cardiol. 2018;4(1):101–113.

Drs. Belhassen and Milman comment

With respect to the present case, we must acknowledge that we have not previously encountered a patient similar to that reported by Drs. Sichrovsky and Mittal.1 A thorough Medline search also confirmed the exceptional nature of this case report.

Based on our latest experience with the Survey on Arrhythmic Events (AEs) in BrS (SABRUS),2,3 we would like to highlight a few points. The SABRUS gathered 678 patients with BrS and AEs from 23 large centers with prior experience in treating BrS. Fifty-nine (8.7%) of the SABRUS patients were female. Although the patient described in the present case is male in all his being, his genetics are of female origin and so we would like to highlight a few critical points in this regard.

Arrhythmic events in “females” with Brugada syndrome

As emphasized by the authors, BrS occurs eight to 10 times more frequently in males despite the disease being equally inherited by both genders. The male/female ratio observed in SABRUS was 10.4; however, this ratio varied according to patients’ age and ethnic origin. Considering the 63-year-old Caucasian, genetically female patient at hand yields a male/female ratio of 2.3 It is noteworthy...
that this ratio is much higher in Asian patients of the same age group (ratio of 14).3

Age at onset of initial arrhythmic event

Adult female patients (identified as those aged older than 16 years) included in SABRUS suffered their first AE at a mean age of 49.5 years ± 14.4 years, which was significantly older than that in males (43.0 ± 12.7 years; p = 0.001).2 We hypothesized that AE onset in females might correlate with circulating estrogen levels because of a predominance for AEs to occur at ages at which estrogen levels are at their lowest.2,3 In the present case, the occurrence of the AE after the age of 60 years could relate more to the marked decrease in estradiol activity rather than to the 20-year use of testosterone.

Mode of arrhythmic event documentation

When considering the patient’s female genetics, it is not surprising that his primary presentation was of an aborted cardiac arrest, as a majority of females in SABRUS (36/59; 61%) developed their initial AE as a first manifestation of the disease.3

Electrocardiogram at the time of arrhythmic event

Similar to our SABRUS female patients (35/59; 59%),3 the ECG at the time of AE did not show a typical type 1 BrS ECG pattern. Interestingly, the latter manifested five months before the event, when the patient was evaluated for an episode of atrial fibrillation.

Quinidine treatment

The authors used a daily low dose of quinidine (quinidine gluconate ER 324 mg), which was similar to the doses of quinidine (≤ 600 mg daily) reported by Marquez et al.4 as effective in preventing AEs in 11 of 14 patients with BrS. It is interesting that such a low dose was apparently effective in both suppressing arrhythmic storms and preventing further AEs during follow-up. This dose is lower than those found to be effective by Anguera et al.5 in 19 (66%) of their 29 patients with multiple AEs (quinidine bisulfate (mean dose: 591 ± 239 mg/day) and hydroquinidine (mean dose: 697 ± 318 mg/day)). In addition, the small dose used in the present paper by Sichrovsky and Mittal3 was much lower than the doses necessary to achieve drug efficacy based on serial electrophysiologic testing6 (ie, mean dose of 1,406 ± 242 mg of quinidine bisulfate and mean dose of 900 mg of hydroquinidine).

Bearing in mind the present outstanding case report, it seems advisable to closely follow with serial ECGs those subjects using chronic exogenous testosterone supplementation. Although anabolic steroids are known to cause sudden death through ischemic coronary disease,7,8 the present case report by Sichrovsky and Mittal highlights the need to consider other possible etiologies such as BrS.
In the fascinating case here reported by Sichrovsky and Mittal, a genetically female patient who presumably was carrying a BrS susceptibility gene variant but who was totally asymptomatic and protected because of her birth gender lost that protection when chronically receiving 200 mg testosterone intramuscularly twice weekly in order to maintain her transgender identity as a male. After experiencing two aborted cardiac arrests at night, the patient was implanted with a dual-chamber implantable cardioverter-defibrillator capable of atrial pacing to avoid the nocturnal bradycardia, which is known to precipitate episodes of ventricular tachycardia/VF. After receiving four appropriate shocks for VF, the patient was placed on quinidine sulfate, which was subsequently changed to quinidine gluconate ER. After a few days, the patient developed angioedema with every dose of quinidine. The addition of the antihistamine cetirizine subsequently allowed him to tolerate once-daily dosing of quinidine. Although quinidine was effective in suppressing all sustained arrhythmias, a high burden of premature ventricular contractions (PVCs) persisted. The PVCs were successfully eliminated via ablation of the endocardial aspect of the RVOT.

Quinidine’s action to suppress ventricular tachycardia/VF in this case is likely attributable to its inhibition of \( I_{io} \). The persistence of the PVCs may be due to the relatively low dose of the quinidine prescribed, owing to a desire to minimize angioedema. In addition to its action to block \( I_{io} \), quinidine also exerts an anticholinergic effect and is an effective inhibitor of the delayed rectifier potassium current \( (I_h) \), which prolongs the QT interval and is the basis for its use in the treatment of short QT syndrome. Both of these actions may have contributed to its ameliorative action in this patient, particularly because, prior to the administration of quinidine, his QT interval was relatively short (QT interval = 300 ms; corrected QT interval = 335 ms) (see Sichrovsky et al.’s Figure 2). Genetic screening was not performed in this case, but the relatively short QT interval suggests the possibility that the case involved a mutation in a calcium channel gene (eg, CACNA1C, CACNB2, or CACNA2D1).

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References

1. Benito B, Sarkozy A, Mont L, et al. Gender differences in clinical manifestations of Brugada syndrome. J Am Coll Cardiol. 2008;52(19):1567–1573.

2. Antzelevitch C. Androgens and male predominance of the Brugada syndrome phenotype. Pacing Clin Electrophysiol. 2003;26(7 Pt 1):1429–1431.

3. Milman A, Andorin A, Gourraud JB, et al. Profile of Brugada syndrome patients presenting with their first documented arrhythmic event: data from the Survey on Arrhythmic Events in Brugada Syndrome (SABRUS). Heart Rhythm. 2018;15(5):716–724.

4. Di Diego JM, Cordeiro JM, Goodrow RJ, et al. Ionic and cellular basis for the predominance of the Brugada syndrome phenotype in males. Circulation. 2002;106(15):2004–2011.

5. Shimizu W, Matsuo K, Kokubo Y, et al. Sex hormone and gender difference—role of testosterone on male predominance in Brugada syndrome. J Cardiovasc Electrophysiol. 2007;18(4):415–421.

6. Fish JM, Antzelevitch C. Cellular and ionic basis for the sex-related difference in the manifestation of the Brugada syndrome and progressive conduction disease phenotypes. J Electrocadiol. 2003;36:173–179.

7. Verkerk AO, Wilders R, de Geringel W, Tan HL. Cellular basis of sex disparities in human cardiac electrophysiology. Acta Physiol (Oxf). 2006;187(4):459–477.

8. Ezaki K, Nakagawa M, Taniguchi Y, et al. Gender differences in the ST segment: effect of androgen-deprivation therapy and possible role of testosterone. Circ J. 2010;74(11):2448–2454.

9. Hayashi H, Miyamoto A, Ishida K, et al. Prevalence and QT interval of early repolarization in a hospital-based population. J Arrhythmia. 2010;26(2):127–133.

10. Matsuo K, Akahoshi M, Seto S, Yano K. Disappearance of the Brugada-type electrocardiogram after surgical castration: a role for testosterone and an explanation for the male preponderance. Pacing Clin Electrophysiol. 2003;26(7 Pt 1):1151–1153.

11. Yamaki M, Sato N, Okada M, et al. A case of Brugada syndrome in which diurnal ECG changes were associated with circadian rhythms of sex hormones. Int Heart J. 2009;50(5):669–676.

12. Barajas-Martinez H, Hu D, Urrutia J, et al. Chronic exposure to testosterone increases expression of transient outward current in human induced pluripotent stem cell (hiPSC)-derived cardiomyocytes (CM). Heart Rhythm. 2013;10(11):1741.

13. Antzelevitch CY, Gj wave syndrome: Brugada and early repolarization syndromes. Heart Rhythm. 2015;12(8):1852–1866.

14. Antzelevitch C, Yan GX, Ackerman MJ, et al. J-Wave syndromes expert consensus conference report: Emerging concepts and gaps in knowledge. Heart Rhythm. 2016;13(10):e295–e324.

15. Belhassen B, Glick A, Viskin S. Efficacy of quinidine in high-risk patients with Brugada syndrome. Circulation. 2004;110(13):1731–1737.

16. Belhassen B, Shapiro I, Shoshani D, Paredes A, Miller H, Laniado S. Idiopathic ventricular fibrillation: inducibility and beneficial effects of class I antiarrhythmic agents. Circulation. 1987;75:809–816.

17. Belhassen B, Viskin S, Antzelevitch C. The Brugada syndrome: is an implantable cardioverter defibrillator the only therapeutic option? Pacing Clin Electrophysiol. 2002;25(11):1634–1640.

18. Alings M, Dekker L, Sadee A, Wilde A. Quinidine induced electrocardiographic normalization in two patients with Brugada syndrome. Pacing Clin Electrophysiol. 2001;24(9 Pt 1):1420–1422.

19. Belhassen B, Viskin S. Pharmacologic approach to therapy of Brugada syndrome: quinidine as an alternative to ICD therapy? In: Antzelevitch C, Brugada P, Brugada J, Brugada R, eds. The Brugada Syndrome: From Bench to Bedside. Oxford, UK: Blackwell Futura; 2004:202–211.

20. Viskin S, Wilde AA, Tan HL, Antzelevitch C, Shimizu W, Belhassen B. Empiric quinidine therapy for asymptomatic Brugada syndrome: time for a prospective registry. Heart Rhythm. 2009;6(3):401–404.

21. Belhassen B, Glick A, Viskin S. Excellent long-term reproducibility of the electrophysiologic efficacy of quinidine in patients with idiopathic ventricular fibrillation and Brugada syndrome. Pacing Clin Electrophysiol. 2009;32(3):294–301.

22. Marquez MF, Bonny A, Hernandez-Castillo E, et al. Long-term efficacy of low doses of quinidine on malignant arrhythmias in Brugada syndrome with an implantable cardioverter-defibrillator: a case series and literature review. Heart Rhythm. 2013;9(12):1995–2000.

23. Pellegrino PL, Di BM, Brunetti ND. Quinidine for the management of electrical storm in an old patient with Brugada syndrome and syncope. Acta Cardiol. 2013;68(2):201–203.

24. Probst V, Evin S, Gournay V, et al. Monomorphic ventricular tachycardia due to Brugada syndrome successfully treated by hydroquinidine therapy in a 3-year-old child. J Cardiovasc Electrophysiol. 2006;17(1):97–100.

25. Schweizer PA, Becker R, Katus HA, Thomas D. Successful acute and long-term management of electrical storm in Brugada syndrome using orciprenaline and quinine/quinidine. Clin Res Cardiol. 2010;99(7):467–470.

26. Hasegawa K, Ashihara T, Kimura H, et al. Long-term pharmacological therapy of Brugada syndrome: is J-wave attenuation a marker of drug efficacy? Intern Med. 2014;53(14):1523–1526.

27. Marquez MF, Rivera J, Hermosillo AG, et al. Arrhythmic storm responsive to quinidine in a patient with Brugada syndrome and vasovagal syncope. Pacing Clin Electrophysiol. 2005;28(8):870–873.

28. Viskin S, Antzelevitch C, Marquez MF, Belhassen B. Quinidine: a valuable medication joins the list of ‘endangered species’. Europace. 2007;9(12):1105–1106.

29. Ohgo T, Okamura H, Noda T, et al. Acute and chronic management in patients with Brugada syndrome associated with electrical storm of ventricular fibrillation. Heart Rhythm. 2007;4(6):695–700.

30. Rosso R, Glick A, Gilksen M, et al. Outcome after implantation of cardioverter defibrillator in patients with Brugada syndrome: a multicenter Israeli study (ISRABRU). Isr Med Assoc J. 2008;10(6):435–439.

31. Belhassen B, Rahkovich M, Michowitz Y, Glick A, Viskin S. Management of Brugada syndrome: a 33-year experience using electrophysiologically-guided therapy with class 1A antiarrhythmic drugs. Circ Arrhythm Electrophysiol. 2015;8(6):1393–1402.

32. Sichrovsky TC, Mittal S. Brugada syndrome unmasked by use of testosterone in a transgender male: gender trumps sex as a risk factor. J Innov Cardiac Rhythm Manage. 2019;10(2):3526–3529.

33. Burashnikov E, Pfeiffer R, Barajas-Martinez H, et al. Mutations in the cardiac L-type calcium channel associated J wave syndrome and sudden cardiac death. Heart Rhythm. 2010;7(12):1872–1882.