DKA-induced Brugada phenocopy mimicking STEMI

CASE PRESENTATION
A 47-year-old Caucasian woman with type 1 diabetes presented with epigastric pain and vomiting. She had not been adherent with her diet and insulin therapy for the past 3 weeks. She never had a personal or family history of arrhythmia-related symptoms, ventricular tachycardia or fibrillation (VT/VF) or premature sudden cardiac death (SCD). Examination revealed dry mucosa, tachycardia and epigastric tenderness to palpation. Her ECG showed ST elevations (V1-V3) with associated T wave inversions (figure 1A). A baseline ECG 1 year ago had no abnormalities. Serial troponin I and T were negative, but Creatinine Kinase MB (CKMB) was elevated. Her biochemistry test showed sodium of 118 mM, potassium of 6.7 mM, bicarbonate of 4 mM, anion gap of 40, glucose of 985 mM and beta hydroxyl-butyrate of >45.0 mg/dL. Cardiac catheterisation revealed normal anatomy with all vessels widely patent; left ventricular end diastolic pressure (LVEDP) was 1 mm Hg. With treatment, diabetic ketoacidosis (DKA) resolved after 8 hours and repeat ECG showed all changes had resolved (figure 1B). She was monitored on telemetry without any VT/VF episodes. Serial ECGs were done with resolution of changes. She had no positive studies for inducible VT. The rest of her admission was uneventful.

QUESTION
Which of the following is the best next step in managing this patient?
A. Quinidine therapy.
B. Implantable cardioverter-defibrillator (ICD) placement.
C. SCN5A gene mutation testing.
D. Observation without therapy.
Despite Brugada syndrome (BrS) being a relatively rare genetic channelopathy, Brugada pattern is an ECG phenomenon, now known as Brugada phenocopy (BrP). The ECG by itself is not diagnostic of BrS; other features include a history of ventricular fibrillation, polymorphic VT, a family history of premature SCD or BrS, induction of programmed ventricular stimulation or pharmacological provocative testing. For patients with the pattern on ECG but none of these other features, BrP is considered. Current guidelines recommend no specific treatment for these patients. A meta-analysis of 1546 cases did suggest that some patients may have subsequent events, but the relative risk was not increased with SCN5A mutation and a number of healthy people harbour the mutation. The acute metabolic changes associated with DKA likely precipitated BrP. Implantation of an ICD in an asymptomatic patient without a spontaneous type 1 Brugada electrocardiographic pattern has no benefit.

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