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Citalopram and the KCNE1 D85N variant: a case report on the implications of a genetic modifier

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Background

Prolongation of the QT interval on the electrocardiogram is clinically important due to the association with an increased risk of sudden cardiac death. A long QT interval may be genetically determined (congenital long QT syndrome) or be drug-induced long QT syndrome e.g. caused by pharmaceutical drugs and electrolyte imbalances.

Case summary

In this report, we describe the case of a 54-year-old woman, who presented with syncope. At presentation, the QTc interval was markedly prolonged, and she was admitted for observation under telemetry. The following day the patient had experienced a near-syncope during an episode of 18 s of Torsade de Pointes (TdP). At the time of TdP, the potassium level (3.4 mmol/L) was mildly reduced, and the ECG showed a QTc interval of 640 ms. In spite of correction of hypokalaemia and discontinuation of the possibly LQTS-inducing drug citalopram the QTc duration remained intermittently prolonged. A transthoracic echocardiogram and a recent coronary angiogram were normal. The patient received an implantable cardioverter-defibrillator. Subsequent genetic testing identified a heterozygous KCNE1 p.D85N (c.253G>A) variant, a known QT modifier with a population prevalence of 1.3%.

Discussion

We conclude that the combination of a commonly prescribed antidepressant, discrete hypokalaemia, and a common KCNE1 QT modifier may cause severe QTc prolongation and life-threatening arrhythmia.

Keywords

Acquired long QT syndrome • Case report • Precision medicine • Genetics ventricular arrhythmias • Torsade de Pointes and KCNE1 p.Asp85Asn (c.253G>A)

Introduction

Prolongation of the QTc interval on electrocardiogram (ECG) is associated with an increased risk of ventricular arrhythmias and sudden cardiac death.1 QTc prolongation results from dysfunction of cardiac ion channels; predominantly altered function of cardiac potassium (I_Kr and I_Ks) or cardiac sodium channels (I_Na) may elicit the phenotype of the long QT syndrome.2 In the congenital form of long

Learning points

• Patients with a KCNE1 D85N variant should avoid reversible risk factors for drug-induced long QT syndrome (dILQTS).
• The finding of a KCNE1 D85N variant can be used to determine whether the relatives share the same increased risk of dILQTS.

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QT syndrome (cLQTS), the ion channels are dysfunctional due to inherited mutations but the same ion channels can also be affected by numerous drugs and hypokalaemia. Drug-induced long QT syndrome (dLQTS) is mainly caused by drugs affecting the promiscuous hERG channel. However, recent studies have also identified an additional underlying genetic disposition or susceptibility to dLQTS: ~30% carry a mutation in the KCNQ1, KCNH2, SCN5A, KCNE1, or KCNE2 genes. These patients can be perceived as having a subclinical form of cLQTS, which will only be expressed when the I\(_\text{Kr}\) current is exposed to a blocking agent. The identification of a disease-causing or a susceptibility inducing mutation may be useful in counselling the proband and relatives at possible risk.

### Timeline

| Time                  | Event                                                                 |
|-----------------------|-----------------------------------------------------------------------|
| 10 months prior       | Started citalopram 20 mg/day for depression                           |
| 3 months prior (in hospital for 2 days) | Hospitalised due to dizziness and headache. An MRI identified an ischaemic lesion in the internal capsule. No sequelae |
| Hospitalization, Day 0 | Malignant syncope, QTc interval at admission 490 ms                  |
| In hospital, +1 day   | QTc interval = 640 ms. 18 s of Torsade de Pointes. Patient felt dizzy but remained conscious during the episode |
| In hospital, +3 days  | Antidepressant was switched from citalopram to the non-QT-prolonging antidepressant sertraline |
| In hospital, +7 days  | Normalization of QT-interval, QTc = 450 ms                           |
| In hospital, +9 days  | Implantation of implantable cardioverter-defibrillator                |

### Case presentation

A 54-year-old woman was brought to the emergency department after a syncope at home. The patient had been cleaning the barbecue in the garden when she felt ‘hard heart beats’ and passed out without further warning. The patient had no memory of the incident that had resulted in a mild head trauma.

Her past medical history was remarkable for deep vein thrombosis (~10 years earlier), ankylosing spondylitis, depression, and a recent transient ischaemic attack (TIA). Approximately 5 years earlier the patient had undergone a coronary angiogram following an episode of chest pains. The angiogram had identified no atheromatosis. Cardiovascular risk factors encompassed hypertension, well-controlled hypercholesterolaemia, and depression. The patient had never smoked and did not have diabetes or a family history of heart disease or sudden premature death. The physical examination, including a cardiac and neurological examination, showed no abnormalities.

At admission, the patient was prescribed the following: citalopram 40 mg/day, adalimumab 40 mg subcutaneously/2 weeks (for ankylosing spondylitis), paracetamol p.n. 1 g max 4 times/day, enalapril 10 mg/day, atorvastatin 40 mg/day, and clopidogrel 75 mg/day.

A CT scan of the cerebrum showed no cerebral damage. The ECG showed sinus rhythm and QTc interval +600 ms (figure 1A). Blood tests were largely normal [serum: potassium 3.4 mmol/L (ref 3.5–4.6 mmol/L) and magnesium, 1.01 mmol/L (ref 0.71–0.94 mmol/L)]. Haemoglobin, leucocytes, creatinine, sodium, alanine transaminase, basic phosphatase, and C reactive protein were within normal ranges. Also, troponin I and D-dimer were not elevated. A transthoracic echocardiography was performed, showing no abnormalities.

After 1 day of telemetry, an episode of self-terminating Torsade de Pointes (TdP) lasting 18 s was recorded (figure 1B). The patient was conscious during the episode but reported feeling dizzy and that symptoms were similar to what she had felt before her syncope the previous day. Empirical treatment with 16 mmol intravenous magnesium and 3 g of potassium chloride was immediately given. Serum potassium increased to 3.8 mmol/L on the following day but the QTc interval remained prolonged (640 ms).

The case was conferred with a psychiatrist, and citalopram was tapered under simultaneous rapid titration of sertraline to 100 mg/day. Citalopram was completely discontinued from Day 5. At 1 week from admission, the QTc duration was within normal range (figure 1C). QTc 450 ms, but the QTc duration fluctuated and was found outside normal range at Day 9 (600 ms), before normalization at Day 15 (460 ms). Earlier ECGs were available for evaluation; previous recordings showed QTc between 440 and 580 ms (figure 1D).

At this point, it was considered possible that the patient was afflicted with the congenital long QT syndrome; even though two known triggers of induced QTc prolongation were present, correction of these did not clearly normalize QTc duration. In addition, previous ECG recordings clearly documented intermittent, significant QTc prolongation over an extended time period. On Day 8 treatment with atenolol 25 mg was initialized. On Day 9, an implantable cardioverter-defibrillator (VVI-ICD) was implanted. Genetic testing was ordered and the proband was found to be heterozygote carrier of the KCNNE1 D85N (c.253G>A) variant.

The patient was closely followed the first 6 months after the TdP episode and seen monthly in the out-patient clinic. Currently, the patient is scheduled for follow-up with her cardiologist yearly, her ICD is interrogated in clinic every 6 months and by remote monitoring every 4 months. In 18 months of follow-up, the ICD has not registered any arrhythmias.

### Discussion

We present a patient with syncope, severe QTc prolongation, and Torsades de Pointes ventricular tachycardia during treatment with citalopram. The patient was a heterozygous carrier of a KCNNE1
DBSN (c.253G>A) variant. The gene encodes for the β-subunit of the voltage-gated potassium channels and the DBSN (c.253G>A) variant has been associated with an increase in QTc interval of ~8 ms.6 Also, the variant is over-represented in patients with diLQTS7 and is a known modifier gene in cLQTS, type one.8 Distinguishing between diLQTS and cLQTS is challenging as there is a significant overlap in the phenotype including QTc duration on the ECG.5 One study reported that 40% of patients with diLQTS may have an underlying cLQTS diagnosis.7 When applying the cLQTS score on our patient, she would receive three points in total (two points for TdP and one for syncope). Three points would categorize the patients as having ‘intermediate probability of cLQTS’.1 However, due to the reversible QT-prolongation, which normalized after discontinuing citalopram and correction of mild hypokalaemia, she was considered to have diLQTS.

Citalopram has a half-life of 36 h, and the patient’s QTc interval was within normal range 4 days (Day 7) after substitution of citalopram with sertraline. Also, we found that on Day 9, the QTc

Figure 1 (A) Twelve-lead electrocardiogram on admission showing prolonged QTc interval (QTc = +600 ms). (B) Telemetry print on Day 2 of admission showing Torsade de Pointes. (C) Electrocardiogram examples during admission with severely prolonged QTc intervals. (D) Historical electrocardiograms used in the evaluation of the patient.
interval was prolonged to 600 ms, which suggests that the QT-prolongation observed on the ECGs from this patient was not solely explained by treatment with citalopram. Consequently, we advised the patient to avoid all QTc prolonging drugs in the future. According to 2015 ESC guidelines, implantation of an ICD in cases with drug-related pro-arrhythmia, should rely on an individual evaluation (Class IIa, Level C). Based on the serial ECG findings in our patient, we remained unassured that her risk of future malignant arrhythmias would be sufficiently controlled solely by avoiding QTc prolonging drugs, therefore, an ICD was implanted. This is supported by the only recent study with long-term follow-up on diLQTS patients with ICDs, where they reported that 44% of patients with an ICD following aborted sudden cardiac death due to diLQTS received appropriate shocks during a mean follow-up of 84 ± 55 months. In contrast, among patients with cLQTS, only 18–28% of patients with cLQTS experience appropriate shocks by their ICD. Though device programming may differ between studies and account for some of this observed difference, one may speculate that we at this point only identify the most malignant subgroup of diLQTS patients since their subsequent risk of ICD shocks is relatively high.

Different drug classes can cause diLQTS and TdP; a frequently updated online tool is available for risk assessment by drug type for physicians and patients. Citalopram is listed in the category of highest risk (‘Known Risk of TdP’). Moreover, a Danish study associated citalopram with a 30% increase in risk for out-of-hospital cardiac arrest. This data supported our concern that citalopram may have elicited the arrhythmias observed in this patient. The genetic test result supported the notion that the patient was at increased risk of drug-induced ECG alterations and hereby at risk of malignant arrhythmias.

Figure 1 Continued.
However, it should be noted that the minor allele frequency of KCNE1 D85N (c.253G>A) occurs with an approximate 1.3% percent frequency in the Danish background population. Also, the patient had several other risk factors for diLQTS: female sex, age, pre-citalopram QTc interval in the high-normal range, and a slightly reduced potassium level. The individual contribution of these distinct risk factors is probably impossible to estimate in the clinical setting. At this point genetic testing and counselling should be tailored to the individual. In this family, one could interpret the findings as the proband as carrying a genetic susceptibility to drug-induced arrhythmias and screen family members accordingly (Figure 2).

Supplementary material

Supplementary material is available at European Heart Journal - Case Reports online.

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Slide sets: A fully edited slide set detailing this case and suitable for local presentation is available online as Supplementary data.

Consent: The author’s confirm that written consent for submission and publication of this case report including image(s) and associated text has been obtained from the patient in line with COPE guidance.

Conflict of interest: none declared.

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