Torsades De Pointes Induced by Levofloxacin in an Unknown Long QT Syndrome

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Clinic Case

A 43-year-old woman arrived to the Emergency Room because of a syncope that lasted approximately 30 s. She denied previous headache, chest pain or any other cardiovascular symptoms before this episode of sudden loss of consciousness.

In her previous medical history, she denied syncope, neurological or respiratory diagnosed diseases. She only referred the history of breast implants without complications 5 years ago, she denied any other chronic disease. She denied drug allergy history, history of trauma and any other prior hospitalization. Her family history denies sudden death and ischemic heart diseases.

Three days before, she reported flu-like symptoms and she was started on levofloxacin and ibuprofen, she denied the intake of some other concomitant medication. At her arrival, vital signs were stable and physical examination was unremarkable except by the finding of a right periorbital hematoma. Her initial laboratories were CK-MB 3 UI/L, ultrasensitive cardiac troponin I 0.70 pg/mL, TSH 3.26 μUI/mL, Na 143 mEq/L, K 4.8 mEq/L, Cl 109 mEq/L, Ca 9.6 mg/dL, Mg 2.1 mg/dL, P 3.4 mg/dL, hs-CRP 0.06 mg/dL, Creatinine 0.6 mg/dL. Her initial EKG (Figure 1A) showed prolonged QTc interval (QTc 520 ms) and ventricular premature complex with R/T phenomenon. A transthoracic Echocardiogram showed preserved left ventricular ejection fraction without regional wall motion abnormalities. At the ER the patient had lipothymy, her second EKG is shown in Figure 1B and shows again a long QTc interval (QTc 500 ms) and helical ventricular tachycardia (Torsades de Pointes). The clinical suspicion at that moment was a drug induced long-QT syndrome induced by the current medication (levofloxacin).

She was treated with magnesium sulphate and during her EKG monitoring she had an asymptomatic, hemodynamically stable helical ventricular tachycardia (Torsades de Pointes) that does not required electrical cardioversion (Figure 1C).

QTc returned to normal after five days free of levofloxacin. No more episodes of helical ventricular tachycardia were noted. Her last EKG is shown in Figure 1D. A cardiovascular magnetic resonance was performed and did not show any data suggestive of structural abnormality of the heart or any cardiomyopathy and specifically ruled out myocarditis, arrhythmogenic right ventricular cardiomyopathy, hypertrophic cardiomyopathy and non-compaction ones. Due to the high-risk of cardiac sudden death an implantable cardioverter was placed along with the proper instruction to the patient and her family to avoid any drug that potentially could prolong the QT interval. The patient is asymptomatic and free of syncope and no arrhythmic events have been register by the cardioverter, therefore she had not received any internal electrical therapy at all until today. Her relatives are currently under investigation and genetic analysis is planned for the near future to this family.

Discussion

In patients documented acquired long-QT syndrome is rarely present Torsades de Pointes. Some factors that predispose to QT prolongation and higher risk for this arrhythmia secondary to drug therapy includes: female sex, low left ventricular ejection fraction, older age, left ventricular hypertrophy, ischemia, bradycardia, electrolyte abnormalities and prolonged repolarization [1,2].

Main types of drugs that prolong the QT-interval include antiarrhythmic drugs (disopyramide, dofetilide, ibutilide, procainamide, quinidine, sotalol, bepridil, amiodarone), anti-infective agents (azole antifungals, fluoroquinolones, macrolide antibiotics, antiretrovirals), antiemetic drugs (domperidone, druperidol), antihistamines (astemizole, hydroxyzine, terfenadine), antipsychotic agents (chlorpromazine, haloperidol, mesoridazine, thioridazine, pimozide) and analgesic drugs (methadone). Other causes, not directly related to drugs, which can prolong QT-interval are electrolyte abnormalities (hypokalemia, hypocalcemia or hypomagnesemia), bradycardia, ischemia, stroke and structural heart disease [1-3].

The correct measure of QT-interval is in lead II of a 12-leads EKG from the onset of the QRS complex to the point where T wave ends and its adequate correction to heart rate by Bazett’s formula makes the diagnosis. Bazett’s formula obtains QTc dividing QT-interval by the square root of the RR interval (in ms). QTc is normal if it is less than or equals to 450 ms in men and less than or equals to 470 ms in women [4,5].

Fluoroquinolones prolong the QT interval by blocking the cardiac voltage-gated rapid potassium channels (I钾). This adverse effect can predispose by the interaction with other drugs that may prolong the QT interval and for the dosage of fluoroquinolone. The chemical composition of fluoroquinolones also found as a predisposing factor for the increase in QT interval prolongation. The probability of this adverse drug reaction with levofloxacin is described 1 per million prescriptions. Females are more susceptible than males to QTc prolongation caused by drugs, because of the specific regulation by sex hormones of ion channel expression and function [6].

The management of this adverse reaction includes the identification of any precipitating drug, along with aggressive correction of any metabolic abnormality. Patients with prolonged QT-interval associated with syncope or Torsades de Pointes should be admitted to the Hospital for EKG monitoring during the period of withdraw and washout of the precipitating drug allowing the treatment of concomitant arrhythmias, if indicated. In specific cases, where acute
therapy is necessary, the proper management includes intravenous magnesium, maintenance of serum potassium levels in the upper limit of normal range, isoproterenol infusion, transient transvenous cardiac pacing, and in some cases the treatment with mexiletine hydrochloride it is also indicated [7].

References