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Editorial

The prolongation of the QT interval of the electrocardiogram is known to be associated with ventricular arrhythmias. Although, anti-arrhythmic drugs are known to prolong the QT interval, this is also prolonged by several non-cardiac drugs that are utilized in the clinical setting [1-6]. The QT interval, a measurement of ventricular systole and diastole, is the electrocardiographic expression of the ventricular depolarization and repolarization. The QT interval is measured from the onset of the QRS to the end of the T wave. This task is not always easy to perform since the last portion of the T wave often camouflages with the isoelectric line.

The influx and efflux of sodium, potassium, and calcium through the cell membrane determines the physiologic depolarization and repolarization of the cardiac cell. Normal depolarization involves a rapid influx of sodium across the cell plasma membrane. Repolarization occurs when the efflux of potassium exceeds the declining inflow of sodium and calcium. These ion currents utilize specific channels that are under the influence of multiple factors. The long QT syndrome is characterized by QT interval prolongation and a tendency to develop a potentially lethal ventricular tachycardia [1,2]. Several anti-arrhythmic drugs, as well as some non-cardiac medications, block a specific potassium channel: $I_{Kr}$. In the case of the anti-arrhythmic drugs, inhibition of $I_{Kr}$ channels is the mechanism for the drug’s therapeutic effect. Non-cardiac drugs that cause the long QT syndrome block these channels incidentally.

Although, the long QT syndrome was associated with the use of anti-arrhythmic drugs at the beginning, this prolongation may complicate the use of a large group of non-cardiac medications frequently used in internal medicine. Mostly prescribed by general physicians and internists, these non-cardiac drugs are commonly prescribed agents, namely, antibiotics, anti-histamines and anti-psychotic agents. It has been estimated that up to 3% of all drug prescriptions are for medications that may prolong the QT interval [3]. Recognition of the long QT syndrome due to non-cardiac drugs is therefore of great importance. The pro-arrhythmic potential of non-cardiac medications can be estimated experimentally in cell cultures, in isolated hearts, as well as, in animal models [4-6]. For example, drug concentrations at which $I_{Kr}$ channels are blocked can be determined in cell cultures. However, these models only provide rough estimates of a given drug’s pro-arrhythmic potential and have limited clinical implications [4].

About five decades ago, Jervell and Lange-Nielsen described the congenital syndrome of deafness, long QT and malignant arrhythmias in infancy [7]. A milder, more common, form of the congenital long QT syndrome was subsequently described by Romano and Ward [8,9]. At that time, the pro-arrhythmic effect of quinidine [10] and some non-cardiac medications [11] were also reported. Although for years the congenital and acquired forms of the long QT syndrome were considered to be two distinct entities [12], it is now clear that all forms of the long QT syndrome are caused by dysfunction of ion channels in the membrane of cardiomyocytes. This ion channel dysfunction may be due to mutations in genes that encode the channel proteins in the congenital syndrome, or may result from the effects of pharmacological agents in the acquired disease [13].

Several non-cardiac medications like antibiotics, anti-histamines and anti-psychotic drugs, block potassium channels in myocardial cells and may thus prolong the QT interval triggering a life-threatening ventricular arrhythmia known as torsade de pointes [14-16]. Most patients with drug induced torsade de pointes have additional clinical risk factors that are readily identifiable such as female gender, organic heart disease, hypokalemia and a history of long QT or drug-induced arrhythmias. For patients without these risk factors it is neither practical nor necessary to implement screening measures such as electrocardiography or measuring serum potassium levels before therapy is initiated. However it is important to consider other preventive measures. For example, it should be avoided the concurrent administration of two or more drugs that prolong the QT interval, as well as, the administration of medications that impair the metabolism of a QT-prolonging drug. Preventing torsade de pointes during drug therapy largely depends on the early identification of patients prone to develop this condition. Predictors of increased risk for QT prolongation with non-cardiac drugs have been described [16]. Of these risk factors, mutations in genes encoding ion channels or drug-metabolizing enzymes can only be detected in selected centers. Moreover, known mutations or polymorphisms associated with an increased risk for arrhythmias are present in only 4-15% of patients with drug-
induced torsade de pointes [14]. In the future, genetic testing may be routinely available to identify patients with mutant ion channels or drug-metabolizing enzyme systems [15].

There is an increased incidence of ventricular arrhythmias in the setting of organic heart disease. Patients with structural heart disease, particularly those with marked left ventricular hypertrophy and heart failure, have abnormal repolarizing currents in the ventricle. Because of their reduced “repolarization reserve” [16] they are at increased risk of developing torsade de pointes when treated with medications that prolong repolarization. It was observed that an average of 42% of patients with torsade de pointes due to non-cardiac medications had structural heart disease. About 19% of patients developed torsade de pointes with anti-histamines, 46% with psychiatric drugs and 52% with antibiotics [17,18]. Concomitant administration of two drugs that lead to QT prolongation should be avoided whenever possible, especially if additional risk factors, namely, female gender, heart disease, are present. Although the risk of torsade de pointes is directly related to the degree of QT prolongation, the exact risk of arrhythmia cannot be predicted by the QT interval. Pro-arrhythmia from non-cardiac drugs appears to be a relatively late complication. In contrast to the pro-arrhythmic effects of anti-arrhythmic medications, which usually appear within 48-72 hours from the onset of therapy [19], only 19% of arrhythmias due to non-cardiac drugs given orally will occur within 72 hours from onset of therapy, and 60% of arrhythmias will occur more than a week after the onset of therapy. If combination therapy is necessary, serial electrocardiograms should be obtained.

It is essential to consider some interesting facts in order to perform an adequate diagnosis and proper therapeutic management. First, the list of medications known to cause the long QT syndrome is constantly expanding. Second, the long QT syndrome should always be considered a potential side effect of drug administration. Third, there are several accessible sources, including Web sites that provide updated information on drug interactions and their direct effects on the QT interval. Fourth, the most important preventive measure is avoidance of drug interactions that prolong the QT interval, and the risk of torsade de pointes may be heightened by the direct effect of multiple drugs on the QT interval or by impaired metabolism of a QT prolonging medication by an additional drug [20-24]. To have these facts in mind will lead to simple precautions and preventive measures that can minimize the risk of serious ventricular arrhythmias in patients taking multiple medications.

References


