Strategies to prevent fatal arrhythmias in patients taking antipsychotics

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Concern has grown about the cardiac effects of antipsychotics as more is learned about rare but deadly torsade de pointes. Which patients require an ECG before you prescribe an antipsychotic? And which agents require the greatest caution?

Before approving the antipsychotic agent ziprasidone last year, the Food and Drug Administration required specific safety data on whether the drug might cause the life-threatening arrhythmia known as torsade de pointes.

The FDA’s action, which delayed the drug’s approval for 3 years, underscores growing concern about the risk of cardiovascular effects with the use of antipsychotic and other agents known to prolong the cardiac QT interval. This concern has led to withdrawal of some drugs before reaching the market (e.g., the atypical neuroleptic sertindole), the addition of “black box” warnings in the labeling of some antipsychotics, and withdrawal from the market of antihistamines terfenadine and astemizole and the GI stimulant cisapride.

continued on page 14
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In torsade, the stimulus for the VT moves within the ventricle, changing its shape from beat to beat. This multifocal VT differs from the more common unifocal VT, in which all the QRS complexes appear the same.

**Drug-induced torsade de pointes**

Although the term torsade de pointes was first described in 1966, the drug-induced form of this arrhythmia has been recognized for nearly a century.

**Quinidine**

Around 1920, cardiologists first used quinidine to help restore normal sinus rhythm in patients with atrial fibrillation, most commonly due to rheumatic heart disease. In 1964, Selzer and Wray studied the use of quinidine to convert atrial fibrillation to normal sinus rhythm in more than 200 patients seen during 4 years in a cardiopulmonary clinic. In a subgroup of eight patients, these researchers documented 10 reactions (including five documented episodes of ventricular fibrillation/ventricular flutter) among 36 syncopeal episodes that developed within 1 to 6.5 hours of quinidine administration. Symptoms were nonspecific and included nausea, faintness, and feeling ill. It is now recognized that torsade de pointes was the principal rhythm disturbance in those eight patients. Syncope usually occurs early in treatment and may be found in 5% to 10% of patients taking quinidine.

**TCAs and antipsychotics**

Tricyclic antidepressants (TCAs) and antipsychotics that have quinidine-like properties (e.g., thioridazine) also may be associated with QT interval prolongation and torsade de pointes. In high doses (particularly in overdose), TCAs may induce widening of the QRS complex. Fowler et al reported episodes of VT in five patients taking thioridazine—one of whom died.

Mehtonen et al reported sudden unexpected deaths associated with antipsychotic or antidepressant drugs among 31 women and 18 men in a survey of autopsies performed from 1985 to 1988 in Finland. The authors documented therapeutic use of phenothiazines in all but 3 of the 49 cases. Thioridazine was involved in more than half the deaths. In 15 of the deaths, thioridazine was the only antipsychotic drug taken. Drugs other than thioridazine were documented in only 5 of the 49 sudden cardiac deaths.
The incidence of torsade is unknown, but it is an uncommon cardiac abnormality. In the United States, torsade probably accounts for less than 5% of the 300,000 sudden cardiac deaths that occur each year. Because torsade de pointes is rare, regulatory agencies and clinicians use the QT interval as a surrogate ECG marker for risk of torsade de pointes. Heart rate can affect the QT interval, so various formulae are used to correct the QT interval for heart rate (QTc).

What is the QT interval? In a normal ECG (Figure 2), the P wave derives from right and left atrial electrical depolarization. The pacemaker of the heart is located in the sino-atrial node (SAN) in the superior portion of the right atrium. From the SAN, electrical signals travel down three intra-atrial pathways, activating the right atrium, then travel to the atrio-ventricular node (AVN). Bachmann’s bundle—a fourth atrial pathway—passes from the SAN to depolarize the left atrium. From the AVN, the electrical signal travels through the left and right bundle branches to activate their respective ventricles.

Electrical depolarization of the left and right ventricles produces the QRS complex. Most of the electrical forces making up this complex arise in the left ventricle, which is much larger than the right ventricle.

The electrical circuitry of the heart activates the left and right atria in such a fashion that these chambers eject blood into their respective ventricles just before these chambers contract. Optimal ventricular filling maximizes ventricular ejection of blood (Starling’s law). Ventricular repolarization (JT interval—electrical recovery) follows ventricular depolarization. On the surface ECG, the JT interval consists of an isoelectric event—the ST segment running from the end of the QRS complex to the beginning of the T wave—and the T wave itself (directional electrical recovery).

The QT interval, then, consists of both ventricular depolarization (QRS complex) and ventricular repolarization (JT interval). Ventricular repolarization makes up by far the greater portion of the QT interval.

Correcting the QT interval (QTc) In 1920, Bazett noted that as the heart rate slowed, the QT interval lengthened. From personal and reported observations, he derived an equation called the Bazett formula that corrects (or normalizes) the QT interval to a heart rate of 60 beats/min (QTc). In the Bazett formula, the QTc interval is the measured QT interval divided by the square root of the RR interval (time between sequential QRS complexes—the determinant of heart rate) measured in seconds (QTc = QT/√RR).

The Bazett formula is most widely used to estimate the QTc interval, although at least 20 other formulae have been developed in response to the original’s perceived inadequacies. Bazett’s formula is used in most automated interpretations of the ECG.

Up to age 55, the normal QTc interval ranges from 350 to 430 msec for men and 350 to 450 msec for women, and it tends to increase with age. Most cases of torsade occur when the QT or QTc interval is greater than 500 msec. A QTc interval between 450 and 500 msec is cause for concern; a QTc interval that exceeds 500 msec is cause for alarm.

Factors that cause variations in QTc
Factors that can affect the QTc interval and increase the risk of torsade de pointes include electrolyte imbalances, medication use and overdose, cardiac disease, liver disease,
Strategies to prevent fatal arrhythmias

About 70% of torsade de pointes cases occur in women.\(^1\) Menstrual cycle QTc interval measurements are stable throughout the menstrual cycle if quinidine-like drugs are not given.

Variations were seen, however, when Rodriguez et al studied the effect of IV low-dose ibutilide (an antiarrhythmic agent known to prolong the QT interval) on the QTc intervals of 58 healthy subjects (38 men and 20 women, ages 21 to 40). During 1 month, men were studied once and women studied three times, coincident with the three phases of the menstrual cycle. The greatest increase in QTc intervals measurements occurred in women during the first half of their menstrual cycles.\(^1\)

Age and cardiovascular disease

Two congenital long QT syndromes may be associated with sudden death, mostly in children and young adults:

- The Jervell and Lange-Nielsen syndrome is marked by severe congenital deafness and autosomal recessive inheritance.
- The Romano-Ward syndrome has normal hearing and autosomal dominant inheritance.\(^2\)

Congenital long QT syndrome (LQTS) occurs in about one in 5,000 births and accounts for about 3,000 to 4,000 deaths per year in the United States. Nine percent of pediatric LQTS subjects present with sudden cardiac death. More than 71% of patients will die before age 15 if not treated.

Elderly persons tend to have longer QTc intervals than do younger subjects, even when both groups are free of cardiovascular disease.\(^2\) Also, age-matched subjects with cardiovascular disease tend to have longer QTc intervals than do those free of cardiovascular disease.

### Table 1

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<thead>
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<th>Risk factor</th>
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Circadian patterns The QTc interval varies throughout the 24-hour day, with nocturnal values about 20 msec greater than daytime measurements. These differences are driven by changes in autonomic (sympathetic and parasympathetic) tone.\(^6\) In 20 normal subjects, circadian variability was 76 ± 19 msec (range 35 to 108 msec) from day to night.\(^7\) This circadian variation may be accentuated in patients with cardiovascular disease.

Sex At birth, QTc interval measurements do not vary by sex.\(^6\) At puberty, however, the male QTc interval shortens and remains shorter than its female counterpart by about 20 msec until age 50 to 55, coincident with a decline in male testosterone levels. This sex difference appears to be androgen driven.

Electrolytes Electrolyte disturbances, particularly...
hypokalemia and hypomagnesemia, may contribute to or even cause QT interval prolongation.\textsuperscript{22}

Hypokalemia prolongs the cardiac action potential and may cause early afterdepolarization, leading to torsade.\textsuperscript{21} Low potassium levels reduce the net outward potassium current during phase 3 of the cardiac action potential. Hypomagnesemia may contribute to gross U wave alternans, lengthening the cardiac action potential and setting the stage for torsade.\textsuperscript{24} Various factors may contribute to electrolyte disturbances, including use of diuretics and excessive vomiting and diarrhea. Even postprandial states may induce hypokalemia.

Intensive exercise and agitation may be associated with hypokalemia.\textsuperscript{27} Serum potassium may be lower in severely agitated patients (3.59 mmol/L) than in mildly agitated patients (3.79 mmol/L). The mean QTc interval of psychiatric emergency patients may be prolonged (453 ± 40 msec),\textsuperscript{7} with QTc intervals of psychiatric inpatients longer than those of psychiatric outpatients. Altered potassium states probably explain these observations. Mechanisms that link exercise and agitation with hypokalemia remain to be elucidated.

**Metabolic factors** Drugs may alter phase 3 potassium flow, thereby disrupting the synchrony of action of individual cardiac cells during repolarization. This change may induce early afterdepolarizations and torsade.\textsuperscript{21}

Five percent to 10% of Americans of European descent have genetic profiles that make them poor metabolizers of drugs that are metabolized by the cytochrome P-450 isoenzyme 2D6. The Pfizer Inc. 054 study assessed the potential for metabolic inhibitors such as paroxetine to raise antipsychotic drug levels in these patients and induce QTc interval prolongation.\textsuperscript{26}

In response to FDA concerns about QTc interval prolongation associated with the use of ziprasidone, Pfizer studied the potential for QTc interval prolongation when antipsychotics are given with and without metabolic inhibitors of cytochrome P-450 isoenzymes 2D6 (paroxetine), 3A4 (ketoconazole), and 1A2 (fluvoxamine). The study population of 183 subjects (mean age: men, 37.1 years; women 38.8 years) was three-quarters young men with schizophrenia, in good health otherwise and possessing normal ECGs—i.e., patients with a low risk of developing cardiac arrhythmias.

Over the course of about 1 week, daily doses were escalated to ziprasidone, 160 mg; risperidone, 8 mg and 16 mg; olanzapine, 20 mg; quetiapine, 750 mg; thioridazine, 300 mg; and haloperidol, 15 mg. Thioridazine (35.6 msec) and ziprasidone (20.3 msec) showed the greatest QTc interval increase following drug administration (Figure 3). Co-administration of a metabolic inhibitor did not further prolong the QTc interval for these two drugs.

Of the six drugs studied, only thioridazine and ziprasidone showed QTc interval increases ≥5% compared with baseline measurements.

Co-administration of a metabolic inhibitor caused the greatest increase in QTc intervals for quetiapine (from 14.5 to 19.7 msec). This value closely approached the steady-state
Thioridazine, for example, is indicated only for patients with schizophrenia who fail to show an acceptable response to other antipsychotic drugs. Its use is contraindicated in patients who take:

- fluvoxamine, propranolol, and pindolol;
- any drug that inhibits the cytochrome P-450 2D6 isoenzyme (e.g., fluoxetine, paroxetine);
- agents known to prolong the QTc interval.

Use of thioridazine also is contraindicated in patients known to have reduced levels of the cytochrome P450 2D6 isozyme, as well as in patients with congenital LQTS or a history of cardiac arrhythmias.

Psychiatrists are advised to read the warnings and prescribing information in the labeling of all antipsychotics for potential cardiovascular side effects.

When the psychiatrist receives a report of suspected QTc interval prolongation on a patient’s ECG, the following steps are recommended:

- Obtain another ECG.
- Assess serum potassium, magnesium, calcium, and thyroid hormone levels.

In patients with confirmed QTc interval prolongation, any complaint of palpitations, presyncope, or syncope are grounds for urgent referral to a cardiologist.

### Recommendations

Taking a careful history is key to cardiovascular assessment before prescribing an antipsychotic. An ECG is indicated for patients with:

- Personal or family history of syncope or sudden death;
- Personal history of angina pectoris, myocardial infarction, congestive heart failure, cardiac arrhythmias, hypokalemia, hypomagnesemia, or significant cardiac risk factors.

The relative cardiovascular risks associated with antipsychotic agents are shown in Table 2.

An ECG also is required or strongly recommended before prescribing the antipsychotic drugs most commonly associated with QT prolongation and torsade de pointes—droperidol, haloperidol in large doses IV (commonly ≥ 100 mg/d), mesoridazine, pimozide, and thioridazine.

The FDA has strengthened the warning labels required for these agents, adding “black box” warnings about the risks of prolonged QTc intervals, torsade de pointes, and sudden death for droperidol, mesoridazine, and thioridazine.

### Bottom Line

The magnitude and extent of QTc interval prolongation as a predictor of life-threatening arrhythmias such as torsade de pointes are becoming more clear. To avoid antipsychotic-induced cardiac complications, psychiatrists are urged to become familiar with and to carefully comply with FDA-approved labeling of all antipsychotic medications.
Related resources

- European Society of Cardiology guidelines: www.escardo.org/scinews/Guidelines/Haverkamp.pdf
- Sudden Arrhythmia Death Syndromes Foundation (SADSF): www.sads.org (800) 786-7723.
- Drugs that prolong the QT interval and/or induce torsade de points. Georgetown Center for Education and Research Therapeutics: www.torsades.org

References


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