Mortality rate is high in psychiatric patients versus general population. An important cause of this increased mortality is sudden cardiac death (SCD) as a major side-effect of psychotropic drugs. These SCDs generally result from arrhythmias occurring when the posology is high and may attain a toxic threshold but also at dosages within therapeutic range, in the presence of risk factors. There are three kinds of risk factors: physiological (e.g., low cardiac rate of sportsmen), physiopathological (e.g., hepatic insufficiency, hypothyroidism) and “therapeutic” (due to interactions between psychotropic drugs and other medicines). Association of pharmacological agents may increase the likelihood of SCDs either by (i) a pharmacokinetic mechanism (e.g., increased torsadogenic potential of a psychotropic drug when its destruction and/or elimination are compromised) or (ii) a pharmacodynamical mechanism (e.g., mutual potentiation of proarrhythmic properties of two drugs). In addition, some psychotropic drugs may induce sudden death in cases of pre-existing congenital cardiopathies such as (i) congenital long QT syndrome, predisposing to torsade de pointes that eventually cause syncope and sudden death. (ii) A Brugada syndrome, that may directly cause ventricular fibrillation due to reduced sodium current through Nav1.5 channels. Moreover, psychotropic drugs may be a direct cause of cardiac lesions also leading to SCD. This is the case, for example, of phenothiazines responsible for ischemic coronaropathies and of clozapine that is involved in the occurrence of myocarditis. The aims of this work are to delineate: (i) the risk of SCD related to the use of psychotropic drugs; (ii) mechanisms involved in the occurrence of such SCD; (iii) preventive actions of psychotropic drugs side effects, on the basis of the knowledge of patient-specific risk factors, documented from clinical history, ionic balance, and ECG investigation by the psychiatrist.

Keywords: sudden death, psychotropic drugs, cardiac arrhythmia, risk factor, side-effect

INTRODUCTION
The overall mortality rate in psychiatric patients is two- to five-fold higher than in the general population (Brown, 1997; Harris and Barracough, 1998; Ruschena et al., 1998; Raedler, 2010) and their life expectancy is approximately 20% shorter (Newman and Bland, 1991). Mortality is higher in male than in female patients, especially when they are young (Hansen et al., 1997). This excess in mortality is partly linked to the use of psychotropic drugs, especially anti-depressants and neuroleptics (Frassati et al., 2004; Ravina et al., 2007b; Alvarez and Pahissa, 2010), but also amphetamines (Yim et al., 2008) and amphetamine-like derivatives, such as methylphenidate (Langendijk and Wilde, 2006) or lithium salts (Yap et al., 2009).

Death occurs more frequently in patients with extra-cardiac (hypokalemia, drug interactions) or cardiac risk factors (bradycardia, ischemia, dilated or hypertrophic cardiomyopathies, arrhythmogenic cardiopathies of the right ventricle, or other arrhythmias). Sudden death has indeed been reported in patients whose medical history included depolarization disorders, e.g.: reentry arrhythmia, such as Brugada syndrome. Right branch block with elevation of ST segment, a characteristic feature of Brugada syndrome, has been described during treatment with several psychotropic drugs (Rouleau et al., 2001). Thus, anti-depressants that selectively inhibit serotonin reuptake (SISR), such as fluvoxamine, or psychotropic drugs that decrease Na+ entry via Nav1.5 channels (SCN5A) should not be prescribed to patients with abnormal Nav1.5 channels (Stürmann et al., 2010) as this could worsen ECG alterations. Moreover, lithium salts induce ECG changes close to Brugada syndrome (Darbar et al., 2005). In addition, the medical history of patients with sudden death of cardiac origin includes repolarization disorders (e.g. prolonged QT interval). Indeed, a congenital or acquired long QT syndrome is an important cause of sudden death owing to pre-existing mutations of ion channels (in particular hERG channels), that predispose to fatal ventricular arrhythmias including torsades de pointe (Tdp) potentially resulting in ventricular fibrillation (VF; Killeen, 2009). Voltage-clamp electrophysiological studies have shown that SISR, such as fluoxetine and norfluoxetine, one of its metabolites, selectively block
hERG channels via two different mechanisms: (i) direct channel blockade, and (ii) disruption of channel protein trafficking (Rajamanvi et al., 2006). Recently, Lee et al. (2009) reported that sulpiride, a benzamide neuroleptic drug, can prolong the QT interval. These authors using the patch-camp technique on Purkinje fibers and mammalian cells transfected with hERG, KCNQ1/KCNE1, KCNJ2, and SCN5A cDNA, and rat myocytes overexpressing IcA showed that sulpiride increased the duration of action potentials owing to a significant reduction in IKr while its effects on INa, ICa, IKs, and IK1 currents were not significant. The prolongation of the QT interval by this neuroleptic drug is therefore linked to a blockade of hERG channels.

The results mentioned above clearly suggest that anti-depressants and neuroleptics can affect several ion channels involved in the control of cardiac action potentials, and thus exert a proarrhythmic activity. A better understanding of genetics in this area can help clinicians to prevent life-threatening adverse events. A number of genes including SCN5A, SCN4B, CACNL1AC, KCNH2, KCNQ1, KCNE1, ANK2, ALG10, KCN2, KCNE2, RYR2, KCND3, KCND2, ACE, NOS1AP, CASQ2 and Rad have been associated with arrhythmias, and are good candidates to define a proarrhythmic genetic profile (Drago et al., 2008). Therefore, before prescribing anti-depressants and neuroleptics drugs, clinicians should be aware of known interactions of several genes (e.g. SCN5A, LQT1, LQT2, LQT3, LQT8) with these drugs in order to achieve a safer therapeutic use. Navedo et al. (2010) have shown the presence of a mutated calcium channel (Cav1.2) in Timo-

The objective of the present paper is to review the risk factors of sudden death of cardiac origin associated with the use of psychotropic drugs, the underlying mechanisms, and preventive measures.

SUDDEN DEATHS OF CARDIAC ORIGIN ASSOCIATED WITH PSYCHOTROPIC DRUGS
Sudden death of cardiac origin associated with the use of psychotropic drugs can occur in patients with pre-existing cardiopathies including ischemic, dilated, and hypertrophic cardiopathies. However, it is possible that psychotropic drugs can by themselves induce such pathological conditions.

PSYCHOTROPIC DRUGS AND ISCHEMIC CARDIOPATHIES
Several psychotropic drugs can induce thrombosis or coronary spasm, or even myocardial infarct (MI). Thorogood et al. (1992) observed a 17-fold increase in the risk of fatal MI (relative risk: 16.9, 95% confidence interval: 3.9–72.8) associated with the use of psychotropic drugs. Moreover, several studies highlighted the high incidence of ischemic cardiopathies correlated to the nature of the psychiatric disease. Thus, Jin et al. (2011) compared the risk of ischemic cardiopathy development in middle age and elderly psychotich patients with the general population over a 10-year period. After excluding patients with a history of coronary disease or cerebral vascular accident, 179 patients were included. Of these, 68 had schizophrenia, 42 mood disorders, 38 insanity, and 31 severe depressions. Selected criteria for the analysis included previous medical and pharmacological treatments, physical examination, and the metabolic profile of patients. In order to assess the risk of developing ischemic cardiopathies according to the Framingham Heart Study (Levy et al., 1990), the authors calculated the risk of coronary disease in each patient, and then compared the relative risk associated with each psychiatric diagnosis. Compared to the general population, they found a 79% risk increase in patients with schizophrenia, 72% with depression, 61% with mood disorders related to psychosis, and 11% with dementia.

In addition, the analysis of the medical history and post-mortem findings from 850 patients with schizophrenia who died from 1952 to 2007 found a tremendous increase in the incidence of pathological cardiac conditions, particularly chronic ischemic cardiopathy, MI and dilated cardiomyopathies. These findings may have been at least partly related to the long-term use of phenothiazine neuroleptics by schizophrenic patients (Volkov, 2009). It seems that the increased incidences of ischemic cardiopathies as well as the high morbidity and mortality rate in patients with schizophrenia are linked to treatments with atypical anti-psychotics. Indeed, such treatments are associated with metabolic disorders including weight gain, dyslipidemia, and deregulation of glucose metabolism, which have been suggested to increase the risk of coronary disease and diabetes in this population (Barnett et al., 2009).

A number of studies highlighted the existence of ischemic cardiopathies discovered at necropsy in patients previously treated with psychotropic drugs and who died of sudden cardiac death (Frassati et al., 2004). Recently, Suvisaari et al. (2010) studied the prevalence of coronary disease and MI in patients with psychotic disorders. They searched for anomalies of the cardiac conduction and the role of anti-psychotic drugs. The study was based on a representative national survey of 8,028 Finnish patients aged over 30 years. The diagnosis of coronary disease and MI was obtained from clinical and ECG findings. A number of ECG recordings evidenced large Q waves suggestive of a former MI. They were significantly more frequent in patients with schizophrenia than in patients without a medical history of psychiatric disorders. Overall, these results support the conclusion that ischemic cardiopathies are more frequent in patients with schizophrenia. One explanation could be at least in part that neuroleptic drugs could play a role by inducing metabolic disorders resulting in ischemic cardiopathies.

PSYCHOTROPIC DRUGS AND CONGENITAL CARDIOPATHIES
The incidence of sudden death associated with psychotropic drugs is very high in patients with various types of congenital cardiopathies.

The causative role of the amphetamine-like drug methylphenidate in sudden deaths occurring in teenagers has been highlighted by Langendijk and Wilde (2006) who reported 25 cases of sudden death defined according to the World Health Organization criteria following the use of methylphenidate and other amphetamine derivatives. The majority of victims were less than 18 years of age and the most fatal arrhythmias developed in teenagers with a congenital cardiopathy including the long QT congenital syndrome or a hypertrophic cardiomyopathy.
Lithium salts that are widely used in the treatment of bipolar disorders are also associated with cardiac complications such as myocarditis (Talati et al., 2009), severe bradycardia (Waring, 2007), prolongation of the QT interval (van Noord et al., 2009), and severe ventricular arrhythmias potentially leading to VF and sudden death (Cruchaudet et al., 2002). Recently, a case of Brugada syndrome was reported in a 78-year-old patient with severe depression treated for 25 years with imipramine, amineptine, levomepromazine, and lorazepam. The addition of lithium salts induced a right branch block with elevation of the ST segment, reminiscent of a Brugada syndrome (Figure 1).

Accordingly, the use of psychotropic drugs that interfere with the fast Na\(^+\) current and can cause reentry arrhythmias (Pacher and Kecskeméti, 2004) is contra-indicated in patients with Brugada syndrome (Antzelevitch et al., 2005; Table 1). Indeed, the density of Na\(^+\) channels is reduced in these patients, which results in depressed membrane depolarization of ventricular cardiomyocytes (Brugada et al., 2005, 2009; Sicouri and Antzelevitch, 2008).

A majority of neuroleptics can prolong the QT interval and lead to TDP. This risk is particularly high in those patients with alterations of a structure or regulatory protein. Prolongation of the QT interval can be due to an improper inactivation of: (i) the Na channel Nav 1.5 (persisting Na\(^+\) current through mutated SCN5A, responsible of LQT3) as is seen in the Romano-Ward syndrome characterized by prolonged QT interval and anomalies of T waves potentially leading to TDP; or (ii) the Ca channel Cav 1.2 (persisting Ca channel current responsible for LQT8) as is seen in the Timothy syndrome combining LQT8 and autism. A prolonged QT interval can also result from the loss of activity in the IKs (LQT1, LQT5) or IKr (LQT2, LQT6) channels, or from alterations of regulatory proteins, such as ankyrin B (LQT4) or caveolin 3 (LQT9) or the KvLQT1 protein (Romano Ward and Jervell syndrome). TDP have also been reported in patients treated with psychotropic drugs that block HERG (IKr) channels (acquired long QT syndrome: LQT1 and LQT2).

### MECHANISMS INVOLVED IN SUDDEN DEATHS OF CARDIAC ORIGIN

Three main mechanisms can be involved in sudden deaths of cardiac origin: electrophysiological disturbances, myocardial ischemia, and myocarditis.

### SUDDEN DEATH DUE TO ELECTROPHYSIOLOGICAL DISTURBANCES

The involved disturbances include either depolarization or repolarization disturbances.

Depolarization in ventricular cells depends on the fast Na\(^+\) inward current (Figure 2), the inhibition of which can induce reentry arrhythmias potentially resulting in VF. Disturbances of ventricular depolarization, which reflect delayed intra-ventricular depolarization, can be induced by many drugs that inhibit the entry of Na\(^+\) ions in ventricular cardiomyocytes. These include neuroleptics (Lo Vecchio et al., 1996) and tricyclic anti-depressants (Leonard et al., 1995) when used alone or combined (Wilens et al., 1990). Moreover, tricyclic anti-depressants behave as class I antiarrhythmics, that is to say they can inhibit the fast Na\(^+\) current (Mladosievicova et al., 1996). They can induce reentry arrhythmias (Figure 3) such as isolated or bursts of ventricular extra-systoles (VES), bigeminy, ventricular tachycardia (VT), ventricular flutters, or even VF and, in this case, sudden death.

Sudden deaths of cardiac origin can also be due to disturbances of repolarization. Psychotropic drugs that prolong the QT interval and can induce TDP include neuroleptics (Warner et al.,

| Therapeutic class | International denomination |
|-------------------|---------------------------|
| Tricyclic anti-depressants | Clomipramine |
| Tetracyclic anti-depressants | Maprotiline |
| Inhibitors of serotonin reuptake | Fluoxetine |
| Phenothiazine neuroleptics | Cyamemazine |

Table 1 | Psychotropic drugs contra-indicated in patients with Brugada syndrome (Antzelevitch et al., 2005).

**FIGURE 1 | Brugada syndrome.** A = normal ECG; B = Brugada syndrome with right branch block and elevation of the ST segment.
FIGURE 2 | Development of action potential in the His–Purkinje system: passive ion movements (phase 0, 1, 2, and 3), and then active ionic movements (ATPase-dependent during diastole). Lower trace: correlation with ECG recording.

FIGURE 3 | Reentry arrhythmias. They affect a restricted area (micro-reentry) or a large area (macro-reentry) of the myocardium. When the excitation wave goes down the intra-cardiac system that is blocked by an area in refractory period (A), it travels around and may excite it in a retrograde direction (the abnormal zone meanwhile comes out of its refractory period, see B). Then, the wave having reached the initial blockade point can, this time, travel in the normal direction (anterograde) to neighbor non-refractory fibers and cause a new premature contraction (C). This can originate an excitation wave rotating within a closed circle and that may propagate to adjacent myocardial tissues. NB: An identical phenomenon may occur in cases of myocardial ischemia, with a slowing down of intra-ventricular conduction in the ischemic area. The excitation wave reaches tissue that is no longer in refractory period and can travel in reverse direction, and thus, may cause reentry arrhythmias.

Among neuroleptics, old neuroleptics can be involved. Mehtonen et al. (1991) reported 49 cases of sudden death including 31 female and 18 male patients. Of these, 46 were treated with a therapeutic dose of a phenothiazine including chlorpromazine alone in 15 patients and combined with a tricyclic anti-depressant in 5 patients. The authors concluded that the cause of death was TDP resulting from prolonged QT interval. Recent neuroleptics can also be involved. Prolongation of the QT interval has been described with olanzapine (Olanzapine: New Preparation, 1999). Risperidone has also been incriminated in prolonged QTc (>450 ms) followed by TDP. Ravina et al. (2007a) reported the death of an elderly woman treated with neuroleptics in which an ECG Holter recording has evidenced TDP that degenerated to VF. In addition, Pacher and Kecskemeti (2004) published prolongation of the QT interval in patients treated with atypical neuroleptics such as olanzapine, risperidone, sertindole, ziprasidone, quetiapine, and aripiprazole. The magnitude prolonged QT interval by aripiprazole has been highlighted by the US FDA in 2002 with 30-ms prolongation in 3.7% of patients treated with 30 mg/day versus 6 ms in patients under placebo. The magnitude of QTc prolongation is dose- and drug-dependent, which reflects the variable propensity of psychotropic drugs to interact with ion channels in the heart. Thoridazine and ziprasidone have been more often incriminated in QT prolongation and TDP, and this is due to the blockade of hERG channels, which facilitates early depolarizing post-potentials that may result in TDP (Figure 4) that can degenerate to VF in some instances. TDP can also be linked to the persistence of the Na+ channel (LQT 3) or calcium channel (LQT8).

NB: The early after-depolarizations (EADs) are related to the activation of L-type calcium channels. These EADs are capable of inducing re-excitation and ventricular arrhythmias. The prolongation of this phase may, especially when it is heterogeneous, lead to torsades de points. This prolongation is either congenital (congenital long QT syndrome) or due to various drug intakes (acquired long QT syndrome, i.e., iatrogenic). Their occurrence is mainly favored by bradycardia and/or hypokalemia. Drugs potentially torsadogenic belong to many classes of drugs (see below).

**Tricyclic anti-depressants**

Curtis et al. (2003) retrospectively studied 4,825,345 patients aged over 18 years and identified 50 drugs with a high potential for inducing prolonged QT interval and 26 drugs that decrease the metabolic clearance of other drugs that prolong the QT interval. Among 1.1 million patients (22.8%) treated with a drug prolonging the QT interval, 103,119 (9.4%) were treated with 2 or more TDP-inducing drugs, or with a drug prolonging the QT interval combined with another drug inhibiting its hepatic metabolism. In addition, 7,249 (0.7%) were treated with 3 or more drugs prolonging the QT interval. Overall, 22% of patients were over 65 years of age and 74% were women. Anti-depressants were involved in nearly 50% of patients with prolonged QT interval. One recent finding is that anti-depressants, which specifically inhibit serotonin reuptake (ISRS), can prolong the QT interval (Rajamani et al., 1999). The most common causes include: (i) drugs such as antiarrhythmics, quinine, halofanthrine, neuroleptics, and tricyclic anti-depressants; (ii) ionic disturbances (hypokalemia); (iii) congenital anomalies (Jervell and Lange-Nielsen syndrome, Romano-Ward syndrome).
Table 2 | Psychotropic drugs recognized by the Advisory Council of Arizona to have a real risk of torsades de pointes.

| International denomination | Commercial denomination | Therapeutic class |
|----------------------------|-------------------------|-------------------|
| Haloperidol                | Haldol®                 | Anti-psychotic/schizophrenia, agitation |
| Thoridazine                | Mellaril®               | Anti-psychotic/schizophrenia |
| Mesoridazine               | Serentil®               | Anti-psychotic/schizophrenia |
| Chlorpromazine             | Thorazine®              | Anti-psychotic/anti-emetic/schizophrenia/nausea |

Table 3 | Psychotropic drugs implicated in the prolongation of the QT interval and in the genesis of torsades de pointes, but with a lack of sufficient evidence.

| International denomination | Commercial denomination | Therapeutic class |
|----------------------------|-------------------------|-------------------|
| Clozapine                  | Clozali®                | Anti-psychotic/schizophrenia |
| Escitalopram               | Lexapro®                | Anti-depressant/major depression/anxiety disorders |
| Escitalopram               | Cipralex®               | Anti-depressant/major depression/anxiety disorders |
| Paliperidone               | Invega®                 | Anti-psychotic, atypical/schizophrenia |
| Quetiapine                 | Serquel®                | Anti-psychotic/schizophrenia |
| Risperidone                | Risperdal®              | Anti-psychotic/schizophrenia |
| Sertindole                 | Serolect®               | Anti-psychotic, atypical/anxiety, schizophrenia |
| Sertindole                 | Serlect®                | Anti-psychotic, atypical/anxiety, schizophrenia |
| Venlafaxine                | Effexor®                | Anti-depressant/depression |
| Ziprasidone                | Geodon®                 | Anti-psychotic/schizophrenia |

However, the risk of torsades de pointes is low when used at recommended doses and/or in patients without risk factors (e.g., bradycardia, electrolyte disturbances and congenital long QT syndrome, in association with inhibitors of CYP 450 responsible for the metabolism of psychotropic drugs concerned). NB. Classification of Arizona does not include all psychotropic drugs that may prolong the QT interval and cause torsades de pointes. This is the case, for example, of pimozide. It is necessary therefore to be especially careful and evaluate the risk for non-listed drugs. NB. Actions to be taken when prescribing neuroleptics and anti-depressants to patients with disorders of repolarization and/or treated by antiarrhythmic drugs or other potentially torsadogenic drugs: Use a neuroleptic or an anti-depressant at low torsadogenic potential. Make sure the ion balance, particularly in serum potassium. Discard, if possible, the use of drugs that induce bradycardia. Perform an ECG and an ionic balance before the introduction of torsadogenic psychotropics (Lin et al., 2004; Narang et al., 2010). A cardiology consultation may sometimes be necessary.

Sudden Deaths Due to Myocarditis and Myocardiopathies

Another mechanism of sudden death associated with the use of neuroleptics is the development of myocarditis. One of the first studies to report on the responsibility of neuroleptics in the generation of myocarditis and myocardiopathy was that of Kilian et al. (1999) on clozapine. This Australian study included 8,000 patients from 1993 to 1999 and described structural lesions in 23 patients.
Cardiac sudden death and psychotropic drugs

(20 men and 3 women, mean age: 36 ± 9 years). Among these 23 patients, 5 had cardiomyopathy of which one died after 36 months of treatment. The 18 remaining patients had myocarditis and 5 died (3 of sudden death and 2 of cardiac insufficiency) within the 3 first weeks of treatment. In addition, the risk of myocarditis associated with clozapine has recently been said to be underestimated (Schulte, 2011). Other reports of neuroleptics-associated myocarditis included fluphenazine, haloperidol, and risperidone (Coulter et al., 2001).

We observed experimentally the development of myocarditis with a number of neuroleptics including amisulpride, haloperidol, levomepromazine, olanzapine, risperidone as well as combinations of levomepromazine and haloperidol or risperidone (Belhani et al., 2006). Indeed, global ventricular hypertrophy was found in rabbits treated with olanzapine and chiefly the combinations levomepromazine-haloperidol and levomepromazine–risperidone; necrosis with amisulpride and haloperidol; endocardial fibrosis with levomepromazine; but no marked lesions with risperidone. Overall, these abnormal findings are similar to those seen in toxic myocarditis induced in previously healthy hearts.

**PREVENTION OF QTc PROLONGATION IN PATIENTS TREATED WITH PSYCHOTROPIC DRUGS**

NB: The evaluation of the risk/benefit ratio for drugs that prolong the QT/QTc interval should take into account morbidity/mortality associated with the disease as well as the lack of treatment of the disease.

The prevention of QT interval prolongation is based on a number of parameters, which should be known to reduce the generation of TDPs and sudden death. It is primarily based on ECG recording and the interpretation of results, on blood ion levels, on the torsadogenic potential of the (psychotropic) drug(s) of concern, and on the existence of risk factors.

Risk factors for the prolongation of the QT interval and the generation of TDPs are of extra-cardiac, cardiac, and drug origin. Drug-induced TDPs mainly develop when two or more drugs are combined, which highlights the role of drug interactions. The existence of one or several risk factors requires ECG recording prior to treatment with psychotropic drugs, especially when the intravenous route is to be used (Meyer-Massetti et al., 2010).

**RISK FACTORS OF TDP**

Many risk factors have been shown to be involved in the development of TDP, including gender (women), pre-existing cardiovascular diseases (especially bradycardia), electrolytic disturbances, and congenital long QT syndrome (Letas et al., 2009; Alvarez and Pahissa, 2010). They must be searched for prior to starting treatment with neuroleptics or anti-depressants, and include extra-cardiac and cardiac risk factors.

**Age, and hepatic and renal function of patients**

Cardiac accidents tend to be more frequent in elderly patients because of their “physiological” decrease in hepatic and/or renal function due to senescence (Viege et al., 2009). Overt hepatic and/or renal failure can also be involved as they reduce the biotransformation and/or elimination of drugs. Adaptation of drug dosage may be needed (Haddad and Anderson, 2002).

**Gender**

Testosterone provides some protection to men (Stollberger et al., 2005; Vieweg et al., 2009). The difference in the susceptibility to TDP across gender appears after puberty only, which suggests that sex hormones may contribute. In male humans, testosterone has been shown to reduce the duration of the action potential and hence of the QTc interval. In contrast, estradiol can potentiate the effects of drugs that prolong the QT interval. Sex hormones have been suggested to alter Ca²⁺ and/or K⁺ currents, which could also account for differences observed in the action of drugs on the repolarization of cardiac cells (Coker, 2008).

**Genetic polymorphism**

Cytochromes P450 (CYP450) are a group of enzymes specialized in the metabolism of drugs that play a pivotal role in biotransformation processes involving xenobiotics (Hasler, 1999; Nelson, 1999). CYP450 isoenzymes are involved in the metabolism of most neuroleptics including clozapine, haloperidol, perphenazine, risperidone, thioridazine, aripiprazole, as well as tricyclic antidepressants and ISRS (Wilkinson, 2005). For instance, more than 25% of drugs including anti-depressants and neuroleptics are metabolized by CYP2D6. CYP2D6 is characterized by a wide genetic polymorphism (approx. 80 identified alleles) and interethenic variability. Slow metabolizers with a very low metabolic activity account for 5–10% of the Caucasian population, and ultra-rapid metabolizers from 1 to 10% of the Caucasian population. They can be affected either by more severe adverse effects or therapeutic inefficacy, depending on whether or not the drug of interest should be metabolized to become active. Intermediate (10–15%) and good metabolizers (65–80%) have a fairly similar metabolic activity. Phenotyping and genotyping can be useful tools to identify at-risk patients and lead to a more effective and safer individualized therapy. This is a very important point. Indeed, the activity of clozapine, which is metabolized by CYP2D6, is increased in case of deficiency and in turn the risk of adverse effects induced by clozapine is increased (Jerling et al., 1994; Kang et al., 2000).

**CARDIAC RISK FACTORS**

**Bradycardia**

Whatever its origin (physiological or pathological), bradycardia can prolong the QT interval by delaying cellular repolarization in patients treated with neuroleptics (Dhein et al., 2008). There is also a higher risk in patients treated with bradycardia-inducing drugs such as digitalis and in patients with bradycardia of pathological origin, such as hypothyroidism (Micallef et al., 2011). Among ECG recorded in 688 psychiatric patients treated with psychotropic drugs, QTc > 460 ms was observed in 96 patients (14%; Sadanaga et al., 2004). To elucidate the relationship between heart rate and duration of the QT interval, these authors measured QT and RR intervals by ECG Holter during 24 h in 97 patients. They showed that for a heart rate of 80/min, the duration of the QT interval was in the range of 400 ms, and for a heart rate of 50/min, the duration exceeded 480 ms. This study stresses the heart rate-dependency of the QT interval duration.

**Slowing of supraventricular conduction**

Any slowing and obviously the presence of an auriculoventricular block (AVB) is an important risk factor of TDP (Yiger et al., 2010).
Table 5 | Different types of CYP 450, their substrates, and inhibitors (Prior and Baker, 2003).

| Types of CYP 450 | Neuroleptic drugs substrates of CYP 450 | Drugs inhibiting CYP 450 |
|-----------------|------------------------------------------|--------------------------|
| CYP 450-1A2     | Clozapine, haloperidol, olanzapine        | Ciprofloxacin, enoxacin, fluvoxamine, cimetidine |
| CYP 450-3A4     | Clozapine, haloperidol, pimozide, ziprasidone, aripiprazole | Fluvoxamine, fluoxetine, ciprofloxacin, itraconazole, ketoconazole, erythromycin, clarithromycin, indinavir, ritonavir, delavirdine, efavirenz, methadone, amiodarone, cimetidine, nefazodone, grapefruit juice |
| CYP 450-2D6     | Clozapine, haloperidol, perphenazine, risperidone, thioridazine, aripiprazole | Fluoxetine, paroxetine, cimetidine, quinidine, ritonavir, amiodarone, clomipramine, chlordiazepoxide, methadone |

NB: It is noteworthy that some anti-psychotics are metabolized by several CYP 450. If one of these CYP 450 is inhibited by a drug, the metabolism of the neuroleptic will be provided by another CYP 450 whose activity is not necessarily inhibited by the same drug. This is the case, for example, of clozapine that is metabolized by the CYP 450 1A2, 3A4, and 2D6. If the activity of CYP 450 1A2 is inhibited by ciprofloxacin, enoxacin, fluvoxamine, and cimetidine, the CYP 450 3A4 and 2D6 ensure, at least in part, the metabolism of clozapine.

Table 6 | Drugs that may prolong the QT interval.

| Therapeutic class | Drugs involved |
|------------------|----------------|
| Class Ia antiarrhythmics | Hydroquinidine, disopyramide |
| Class III antiarrhythmics | Amiodarone, sotalol, ibutilide |
| Class IV antiarrhythmics | Bepridil |
| Antibiotics (macrolides) | Erythromycin |
| Synthetic antimalarial agents | Halofantrine |
| Anti leishmania agents and trypanocides | Pentamidine diethionate |
| H1 antihistaminics | Mizolastine: this drug slightly lengthens the QT interval |

Table 7 | Drugs responsible for hypokalemia.

| Hypokalemia and mechanisms for its occurrence | Drugs involved |
|-----------------------------------------------|----------------|
| Prokinetic effect | Cisapride (Prepulsid®) |
| Digestive loss | Laxatives, Kayexalate |
| Urinary loss | Glucocorticoids, Mineralocorticoids, High ceiling diuretics and thiazides, Glycyrrhizinic acid, β-Lactamines at high doses, Aminosides, Amphotericin B (by acute tubular necrosis), β2 Mimetics (IV): salbutamol, adrenaline |
| Transfer of intracellular K+ | Blood alkalinizing drugs |

Channelopathies
The presence of Romano Ward or Timothy syndrome considerably increases the generation of TdP associated with psychotropic drugs that can prolong the QT interval. It is absolutely required to evaluate this risk and adapt treatment.

Drug-related risk factors: Drug interactions
Pharmacokinetic interactions: enzyme inhibitors
Such interactions develop in particular when psychotropic drugs are associated with drugs that interact with various CYP450 (Prior and Baker, 2003). Indeed, the combination of a psychotropic drug with another drug that inhibits the enzyme involved in metabolism can induce an increase in plasma levels, and potentially tissue levels with resulting toxicity. The main psychotropic drugs and CYP450 involved in their metabolism and specific enzyme inhibitors are summarized in Table 5.

Pharmacodynamic interactions
The use of potentially torsadogenic neuroleptics increases the risk of TdP when associated with another drug that can prolong the QT interval (Cubeddu, 2003; Table 6). Such an association should therefore be avoided or considered only very carefully.

Hypokalemia (Frimas et al., 2008)
Hypokalemia of pathological or drug-induced origin can shift potential toward less negative values and thus considerably increase the risk of TdP. It is therefore recommended to avoid the combination of neuroleptics with hypokalemiant drugs or

FIGURE 5 | How to handle prescription of a potentially torsadogenic psychotropic in patients at-risk.
attempts to correct hypokalemia. Hypokalemia can have various origins (Table 7):

CONCLUSION

The cardiovascular toxicity of psychotropic drugs can display various clinical manifestations including orthostatic hypotension, syncope, myocardial ischemia, myocarditis, cardiac arrhythmias (reentry arrhythmias, torsade de points...). Among which some may lead to sudden death. These manifestations, in particular sudden death, are more common in patients with congenital or acquired cardiac risk factors, including hypertrophic and/or dilated cardiopathies, ischemic cardiopathies, combination of prescribed drugs that prolong QRS complexes or QT intervals, or drugs that can induce hypokalemia or bradycardia. Particular precautionary measures should be taken including blood ionic balance, ECG recording, and even cardiac echography, to reduce the risk of cardiac adverse events that may be life-threatening (Figure 5).

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