Predicting Plasma Olanzapine Concentration Following a Change in Dosage: A Population Pharmacokinetic Study

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Key words
- olanzapine
- plasma concentration
- population pharmacokinetics
- therapeutic drug monitoring

Abstract

Introduction: Due to high inter-individual variability in peripheral pharmacokinetic parameters, dosing of antipsychotics currently relies on clinical trial-and-error, and predicting antipsychotic plasma concentrations before changing a dose has been a challenge.

Methods: Patients with schizophrenia receiving a stable dose of olanzapine were included. 2 plasma samples were collected at 2 given time points for the measurement of plasma olanzapine concentrations. At least 7 days after a dosage change of olanzapine, a third sample was collected. The plasma concentration of the third sample was predicted in a blinded fashion using a mixed-effects model with NONMEM®, using the following information: the 2 baseline plasma concentrations, the interval between the last dose and blood draw, and clinical and demographic information.

Results: 31 subjects (mean±SD age = 56.0±11.6; 19 men) were enrolled. The mean prediction concentration was observed between the observed and predicted concentrations of the third sample (r = 0.91, p < 0.001).

Discussion: Plasma olanzapine concentrations following an actual dosage change can be predicted in advance with a high degree of certainty.

Introduction

Antipsychotic drugs, which exert their clinical effects mainly through the blockade of dopamine D2 receptors [1], are widely used for the treatment of schizophrenia across the patient’s life span [2]. Positron emission tomography (PET) studies have identified that optimal clinical response in young patients with schizophrenia is related to a “therapeutic window” of the striatal dopamine D2 receptor occupancy of 65–80% [1, 3, 4]. On the other hand, the clinical application of PET to individualized antipsychotic dosing at the bedside has not been feasible because of its low availability and high cost. To tackle this situation, the authors have developed a model to estimate dopamine D2 receptor occupancy levels from plasma concentrations of various antipsychotics, including olanzapine, with a high degree of accuracy [5]. Therefore, individualized dosing with the measurement of antipsychotic plasma concentrations could become a real potential clinical application; the current dose adjustment still relies on a trial-and-error strategy that carries a risk of relapse and adverse effects.

Given high inter-individual variability in peripheral pharmacokinetic parameters [6], the next hurdle is to accurately predict antipsychotic plasma concentrations before an actual dosage change of antipsychotic drugs for each individual. In contrast to conventional pharmacokinetic modeling, the population pharmacokinetic model can, in theory, predict individual pharmacokinetic parameters for antipsychotics, including peak and trough plasma concentrations, using blood samples collected at any given time points with the information on demographic and clinical characteristics of the subjects and dosing information (i.e., dose, time of last dose, and duration between the last dose and blood draw). Thus, the corresponding peak and trough dopamine D2 receptor occupancy levels could also be estimated using the above-mentioned prediction model [5, 7]. In theory, this model could be used to find doses which would result in estimated peak and trough drug concentrations corresponding to the established therapeutic window.
of dopamine D\textsubscript{2} receptor occupancy in the treatment of schizophrenia [8]. The authors have successfully reported on the validity and reliability of predicting plasma concentrations of risperidone and 9-hydroxyrisperidone associated with a dosage change, using population pharmacokinetic techniques [9].

Olanzapine is one of the most frequently and widely used antipsychotic drugs [10, 11]. This drug is characterized by its effect on not only psychotic but also anxiety and depressive symptoms in patients with schizophrenia [12, 13]. Unfortunately, the use of olanzapine occasionally results in a variety of side effects, including cardiovascular and motor side effects as well as impairment in subjective well-being, some of which are at least in part dose-dependent [3, 14, 15]. Moreover, this drug may increase the risk of weight gain and diabetes mellitus in a dose-dependent fashion [16, 17], although this dose-dependence has not always been a consistent finding in the literature [18, 19]. These findings clearly emphasize the need for using the lowest possible dose of olanzapine in the treatment of schizophrenia.

As described above, use of the population pharmacokinetic model and the dopamine D\textsubscript{2} receptor occupancy estimation model could bring psychiatric practice closer to the goals of personalized medicine: enabling the establishment of individualized antipsychotic doses that would result in optimal D\textsubscript{2} receptor occupancy and clinical response.

In this study, the authors used the collective data from 3 clinical trials and tested the validity and reliability of predicting plasma concentrations of olanzapine following a dosage change from baseline concentrations, using population pharmacokinetic techniques [20]. The predicted post-adjustment concentrations of olanzapine were compared to the actually measured concentrations in order to determine the reliability of our prediction strategy.

Methods

Subjects and settings

Data used for the present analysis were derived from 3 clinical trials; in those 3 trials, plasma olanzapine concentrations were measured twice before a change in dosage and once after the change. The first study was designed specifically for the present analysis. In this first study, male or female patients of any race who fulfilled the following criteria were included: (1) diagnosis of schizophrenia, schizoaffective disorder, schizophreniform disorder, delusional disorder, or psychotic disorder not otherwise specified according to the Diagnostic and Statistical Manual of Mental Disorders, Fourth Edition [21], (2) having changed a dose due to insufficient treatment response or side effect, and (3) having been treated with oral tablet form of olanzapine at a steady dose for at least 7 days. The study was conducted at Keio University Hospital, Minami-Hannou Hospital, and Asakadai Mental Clinic and was approved by the institutional review board at all participating sites. The study subjects provided written informed consent after receiving detailed information about the protocol. The trial was registered at the University Hospital Medical Information Network (UMIN) Clinical Trials Registry as UMIN000002294. The second study was an open-label, 28-week, randomized controlled trial to investigate the impact of olanzapine or risperidone dose reduction by half on cognitive function in stable patients with schizophrenia. In this study, clinically stable patients with schizophrenia receiving a stable dose of either olanzapine > 5 mg/day or risperidone > 2 mg/day as antipsychotic monotherapy for at least 3 months were included [22] (UMIN Clinical Trials Registry, UMIN000001834). In the third study, patients aged 50 and older with schizophrenia spectrum disorders who had been treated with olanzapine at a steady dose of ≥ 10 mg/day or risperidone at a steady dose of ≥ 2 mg/day for 6–12 months were included [23] (ClinicalTrials.gov, NCT00716755). This study included PET scanning to measure dopamine D\textsubscript{2} receptor occupancy before and after a gradual dose reduction of antipsychotics by up to 40%.

Study description

The design of the present analysis is summarized in Fig. 1. 2 plasma samples were taken at 2 different random time points for the measurement of plasma olanzapine concentrations. At least 7 days after a dosage change of olanzapine, the third sample was collected at any given time point.

The plasma olanzapine concentration of the third sample was kept blind to the investigator (R.R.B.), who calculated the predicted plasma concentrations using NONMEM VII [24]. Plasma concentrations of olanzapine were assayed by hiruphenized plasma using liquid chromatography with tandem mass spectrometry at the Centre for Addiction and Mental Health, Toronto, Canada. The following information was also collected: concomitant medications, intervals between last dose and blood draw, and demographic and clinical information, including age, diagnosis, sex, race, and smoking status. All subjects recorded the time of their last dose before the scheduled collection of blood samples, and concomitant medications were kept constant throughout the study period. In all subjects, medication adherence was evaluated by their verbal report. We excluded the subjects who lacked any information needed for population pharmacokinetic analysis (e.g., dosing interval, weight).

Population pharmacokinetic analysis

Using a previously established population pharmacokinetic model for olanzapine [20], the plasma concentration of olanzapine at the time of the third blood draw (i.e., the blood draw after a dosage change) was individually predicted with the 2 plasma concentrations of olanzapine at the baseline dose, interval between last dose and blood draw, age, and smoking status (Fig. 1). The mixed-effects model was established using the data derived from the Clinical Antipsychotic Trials in Interven-
tion Effectiveness (CATIE). This dataset included olanzapine concentrations (1527 plasma samples) from 523 subjects (age range, 18–103 years); a one-compartment mixture model with additive error best described the data [20]. According to the results of this trial, sex, race, and smoking habits that were found to affect olanzapine clearance were included in this population pharmacokinetic model for the present study.

Statistical analysis
Statistical analyses were carried out using the IBM SPSS Statistics Version 21 (IBM Corporation, Armonk, NY, USA) and the GraphPad Prism Version 5 (GraphPad Software, Inc, San Diego, CA, USA). The predictive performance of this procedure was assessed with the mean prediction error and the root-mean-squared prediction error [25]. The prediction error refers to the difference between the true (i.e., measured) values and those predicted by the model. The mean squared prediction error is defined as a mean of squared values of prediction errors for all data points. The root-mean-squared prediction error refers to the root of the mean squared prediction error, which is a measure of “bias.” Pearson correlation analysis was also used to examine the relationship between the observed and predicted values although this is a less precise value of predictive performance. A p-value of <0.05 was considered statistically significant (2-tailed).

Results
Data from a total of 31 subjects (3 outpatients from the first study, 2 inpatients and 7 outpatients from the second study, and 19 outpatients from the third study) were used for the present analysis; their demographic and clinical characteristics are summarized in Table 1. The dose of olanzapine was increased and decreased in 2 and 29 subjects, respectively (mean ± SD dose, mg/day: 5.0 ± 0 to 7.5 ± 0 and 18.1 ± 7.7 to 11.1 ± 5.3, respectively). Inter-individual variations in the oral clearance of olanzapine were as wide as approximately 4-fold (range, 12–47l/h). The mean (95% confidence interval) prediction errors and root-mean-squared prediction errors (ng/mL) were as low as 1.6 (−2.8 to 6.0) and 12.0 (8.9 to 15.0), respectively (Fig. 2). The observed and predicted concentrations of olanzapine were highly correlated (r = 0.91, P < 0.0001) (Fig. 2). The distribution of the prediction errors is shown in Fig. 3.

Discussion
To the best of our knowledge, this is the first attempt to predict the plasma concentration of olanzapine following a dosage change in patients with schizophrenia spectrum disorders with the use of population pharmacokinetic techniques. We have previously reported that plasma concentrations of risperidone can be reliably predicted prior to a dose adjustment using the same population pharmacokinetic technique [9]. Consistent with our previous report, the present study demonstrates that the same methodology can also be applied to olanzapine in a reliable manner. Considering that a model to estimate the dopamine D2 receptor occupancy from plasma concentrations of olanzapine is available [4], the results of this study can be used to predict peak and trough D2 receptor occupancy levels before dose adjustment of olanzapine in individual patients without performing a brain imaging study. This is of high clinical pertinence for the treatment of schizophrenia, since excessive striatal D2 receptor blockade has consistently been associated with extrapyramidal side effects [1,3,4]. Furthermore, excessive dopamine D2 receptor blockade also increases the risk of cognitive impairment [26] and hyperprolactinemia [27]. On the other hand, insufficient D2 receptor blockade has been shown to be associated with poor clinical response in the acute phase of the illness [1,3]. These findings emphasize the importance of targeting an appropriate dopamine D2 receptor blockade of 65–80%, the so-called “therapeutic window”, in the acute phase of treatment for persons

Table 1 Demographic and Clinical Characteristics of Subjects (N = 31).

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>Values</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age, mean ± SD (range), years</td>
<td>56.0 ± 11.6 (30–72)</td>
</tr>
<tr>
<td>Male, N (%)</td>
<td>19 (61.3 %)</td>
</tr>
<tr>
<td>Smoking, N (%)</td>
<td>10 (32.3 %)</td>
</tr>
<tr>
<td>Race</td>
<td></td>
</tr>
<tr>
<td>Caucasian, N (%)</td>
<td>27 (87.1 %)</td>
</tr>
<tr>
<td>African, N (%)</td>
<td>4 (12.9 %)</td>
</tr>
<tr>
<td>Inpatient, N (%)</td>
<td>7 (22.6 %)</td>
</tr>
<tr>
<td>Dose of olanzapine, mean ± SD (range), mg</td>
<td>17.2 ± 8.1 (3.8–35.0)</td>
</tr>
<tr>
<td>Before dosage change</td>
<td></td>
</tr>
<tr>
<td>After dosage change</td>
<td>10.9 ± 5.2 (2.5–22.5)</td>
</tr>
</tbody>
</table>

![Fig. 2](image-url) Relationship between observed and predicted plasma concentrations of olanzapine, using population pharmacokinetic techniques.

![Fig. 3](image-url) Distribution of prediction errors.
with schizophrenia. Moreover, physicians as well as patients are often reluctant to reduce the antipsychotic dose during the maintenance treatment of schizophrenia in fear of potential relapse. As a result, patients continue to be maintained on higher doses of antipsychotics than actually needed. If the lowest possible effective dose can be estimated for each individual, such reluctance will be dispelled and the situation will change for more individualized treatment.

Using widely available drug assays for 2 blood samples collected at any given time point, the population pharmacokinetic model could be used to determine a new dose resulting in estimated peak and trough drug concentrations that would correspond to the therapeutic window of dopamine D2 receptor occupancy. According to the model for the estimation of dopamine D2 receptor blockade from plasma antipsychotic concentrations, the therapeutic window of 65–80% corresponds to plasma concentrations of olanzapine of 13.9–43.6 ng/mL [5]. Similarly, when antipsychotic treatment is initiated, the minimal effective oral dose of olanzapine associated with the targeted dopamine D2 receptor occupancy could also be individually estimated. This can be accomplished by measuring the baseline plasma drug concentrations at steady state of a starting test dose of olanzapine, such as 2.5 mg oral dose per day for 1 week. Thus, personalizing antipsychotic doses to ensure optimal clinical response based on estimated striatal D2 receptor occupancy may one day become common clinical practice; using plasma drug concentrations and population pharmacokinetic models [7].

While the utility of therapeutic drug monitoring (TDM) of olanzapine for the treatment of schizophrenia has still been controversial [28], a number of studies have suggested a link between its blood concentration and clinical outcomes [29–32]. For example, Fellows et al. identified a maximal plasma olanzapine concentration of 23 ng/mL as a cut-off point to distinguish responders from non-responders [29]. Moreover, according to Mauri et al., plasma olanzapine concentrations ranging from 5 to 120 ng/mL showed a curvilinear correlation with clinical improvement in patients with schizophrenia [31]. Furthermore, using a large naturalistic dataset from an olanzapine TDM service, Patel and colleagues found that for a given dose of olanzapine female non-smokers had higher plasma concentrations than male smokers. This data suggests that TDM of olanzapine may be useful not only in assessing adherence, but also in limiting oral doses to reduce the risk of long-term toxicity [32]. In 2011, the Arbeitsgemeinschaft für Neuropsychopharmakologie und Pharmakopsychiatrie (AGNP) released the consensus guidelines for TDM of olanzapine, suggesting an optimal therapeutic range of plasma olanzapine concentrations between 20–80 ng/mL [33]. Notwithstanding such recommendations, TDM is rarely performed for olanzapine in clinical practice. Law et al. recently conducted an intervention study to test the clinical utility of TDM for olanzapine. They found that TDM could be feasibly implemented, potentially aiding dose optimization, supporting the routine clinical use of TDM-based dose adjustment for olanzapine [34]. To summarize, the evidence of TDM for olanzapine thus far indicates that measuring the plasma concentration of olanzapine seems to have the potential to guide physicians in determining its effective dose, especially in the face of insufficient response and dose-dependent side effects.

There are some limitations to the present study worth noting. First, although the predictive performance was assessed with the mean prediction error and the root-mean-squared prediction error [25], there was no other model to compare the accuracy of the results. Second, if we sought for the individualized treatment with antipsychotics, it might have been ideal to include genetic information. Third, our conclusions cannot be generalized to antipsychotics other than olanzapine and risperidone, although population pharmacokinetic models have been established for other antipsychotics, including ziprasidone [35], clozapine [6], and perphenazine [36]. Fourth, the pharmacokinetics of olanzapine follows linear and dose-proportional effect within the approved dosage range [37]. However, it may be nonlinear and disproportionate beyond a certain plasma concentration of olanzapine. Fifth, since this sample size was small, the statistical power may not be sufficient to draw conclusions. The findings in the present study should be confirmed by future investigations with larger sample sizes. Finally, our proposed model-guided antipsychotic dosing still requires further validation. Therefore, we have been performing a series of clinical trials to test the reliability of our prediction model in estimating target doses in the treatment of schizophrenia.

In conclusion, the results of the present study demonstrate that plasma olanzapine concentrations can be reliably predicted prior to an actual change in dosage using population pharmacokinetic techniques. Considering the close relationship between plasma concentrations, dopamine D2 receptor occupancy, and clinical (psychotherapeutic and adverse) response, these findings suggest that individualized dosing with the measurement of antipsychotic plasma concentrations, instead of the conventional trial-and-error strategy, may become a clinical option for the treatment of schizophrenia.

Author Contributions

Dr. Uchida led the 1st study. Dr. Takeuchi led the 2nd study. Dr. Mamo and Graff-Guerrero led the 3rd study. Drs. Bies and Uchida designed the study. Drs. Tsuboi, Bies, and Uchida analyzed the data. Drs. Tsuboi and Uchida wrote the first draft of the manuscript. All authors have contributed to and approved the final manuscript.

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Previous Presentation

Some of the data were presented at the 23rd meeting of the Japanese Society of Clinical Neuropsychopharmacology, Okinawa, October 24, 2013.

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Conflicts of Interest and Financial Disclosure

Dr. Tsuboi has received manuscript fees from Dainippon Sumitomo Pharma and speaker’s honoraria from Eli Lilly, Tsumura, Yoshitomi Yakuhin, Dainippon Sumitomo Pharma, Kracie Pharma and Mitsubishi Tanabe Pharma within the past 2 years. Dr. Bies has received NIH, CAMH, Lilly and Indiana University grant funding.

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Dr. Pollock receives research support from the National Institute of Health and the Canadian Institutes of Health Research. Within the past 5 years, he has been a member of the advisory board of Lundbeck Canada (final meeting was May 2009) and Forest Laboratories (final meeting was March 2008). Dr. Pollock has served one time as a consultant for Wyeth (October 2008). He was also a faculty member of the Lundbeck International Neuroscience Foundation (LINF) (final meeting was April 2010).

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References

8 Nakajima S, Uchida H, Bies RR et al. Dopamine D2/3 receptor occupancy following dose reduction is predictable with minimal plasma antipsychotic concentrations: an open-label clinical trial. Schizophrenia Bull 2015; (in press)
18 Mitchell M, Riesenberg R, Bari MA et al. A double-blind, randomized trial to evaluate the pharmacokinetics and tolerability of 30 or 40 mg/d oral olanzapine relative to 20 mg/d oral olanzapine in stable psychiatric subjects. Clin Ther 2006; 28: 881–892

26 Sakurai H, Bies RR, Stroup ST et al. Dopamine D2 receptor occupancy and cognition in schizophrenia: analysis of the CATIE data. Schizophr Bull 2013; 39: 564–574
29 Fellows L, Ahmad F, Castle DJ et al. Investigation of target plasma concentration-effect relationships for olanzapine in schizophrenia. Ther Drug Monit 2003; 25: 682–689