Bioequivalence Study of Two 50 mg Desvenlafaxine Extended Release Formulations: A Randomized, Single-Dose, Open-Label, Two Periods, Crossover Study

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Abstract

This is a pharmacokinetic study of two formulations containing Desvenlafaxine succinate 50 mg extended release. Its objective was to compare the bioavailability between the Test product (Desvenlafaxine ER produced by Tecoquimicas S.A., Colombia laboratory) and the Reference product (Pristiq XR® produced by Wyeth laboratory) and to be able to determine the Bioequivalence between the both of them. For this, an open label, two periods, two previously randomized sequences, crossover, single postprandial 100 mg dose study with an 8 days washout period between each period in 24 healthy volunteers was performed, including the collection of 13 plasma samples within 0 and 48 hours from all volunteers who participated in the clinical phase. The analytical method used was High Performance Liquid Chromatography (HPLC) with UV detector. The obtained mean peak concentration (Cmax) for the Test and Reference products were 215.8 ng/mL and 196.9 ng/mL and for the area under the curve up to 48 hours (AUC0-t) 3849,6 ng.h/mL and 3605,4 ng.h/mL respectively. The 90% confidence interval for the Cmax parameter is within the range of 103,58-113,63 and for the AUC0-t parameter, the 90% confidence interval is within 97,96-111,39. Based on the FDA, EMA and WHO bioequivalence investigation guidelines, the CI is within the allowed ranges for bioequivalence and interchangeability declaration of the Tecoquimicas S.A. product with the Reference product.

Keywords: Bioequivalence; Desvenlafaxine; Antidepressant; Pharmacokinetics

Introduction

Desvenlafaxine is the O-desmethyl active metabolite of venlafaxine, which presents a dual mechanism of action inhibiting the serotonin and adrenaline recapture in the synaptic cleft through serotonin (SERT) and noradrenaline transporters (NERT) blocking in the presynaptic membrane. It was approved in 2008 by the FDA for the Major Depressive Disorder (MDD) treatment given its short-term safety, tolerability and demonstrated efficacy in placebo-controlled trials [1-3]. In its succinate salt form, an extended active ingredient release is produced, allowing the administration of one tablet every 24 hours since it has a plasma concentration peak at 6 to 8 hours in fasting conditions (8 to 10 hours if taken postprandial) and an elimination half-life between 9 to 15 hours [4,5]. It has linear kinetics in the 100 to 600 mg/day range, although its antidepressant efficacy has been demonstrated with 50 mg/day doses. Its oral Bioavailability is 80% and it is not modified by low fat meals [4,5]. It is mainly metabolized by glucuronidation through UDP-glucuronyltransferase and to a lesser extent through the cytochrome CYP3A4 oxidative pathway. This pharmacokinetic characteristic decreases its interindividual variability given the slightest polymorphism observed in this enzyme, opposite to the metabolism description of venlafaxine, which is mainly metabolized by the CYP2D6 cytochrome pathway, an enzyme likely to present such polymorphisms.

The major depressive disorder is a mental disease with a high prevalence in adult population (close to 7%) and a prediction of becoming the main world disability cause by 2030 [5,6]. This pathology has an adverse impact in patients with related diseases, such as cardiovascular disease and chronic pain handling, being the pharmacological treatment the main support for its treatment.

The objective of this study was to establish the Bioequivalence of two formulations containing Desvenlafaxine succinate 50 mg tablets by comparing its Bioavailability after a single dose between the Test product (produced by Tecoquimicas S.A., Colombia) and the Reference product (Pristiq XR® produced by Wyeth).

Materials and Methods

Study formulations

The Test formulation was Desvenlafaxine 50 mg extended release tablets produced by Tecoquimicas S.A, Cali-Colombia (Batch EX-3D09) and the Reference product was Pristiq®-Desvenlafaxine 50 mg extended release tablets produced by Wyeth (Batch 82386).

Physical-chemical properties, active ingredient assessment, dose uniformity and dissolution profile were evaluated for both the Test and Reference products in order to determine the Pharmaceutical Equivalence of these before carrying out the in vivo study.

Subjects

Twenty four healthy non-smoking female and male subjects, aged between 18 and 42 years old with a Body Mass Index (BMI) of
19-25 kg/m² participated. All volunteers were assessed with a medical examination and laboratory tests before the clinical phase initiation to confirm their health condition. Alcoholism history, preexistent diseases compromising liver or kidney function, blood dyscrasia or proteinuria were considered as exclusion factors.

**Medical examinations and clinical laboratory tests**

Performed clinical laboratory tests included CBC, total and direct bilirubin, creatinine, glycemia, total protein, complete urinalysis, HIV Elisa test, antibodies against hepatitis B and C, electrocardiogram and blood pregnancy test for women.

**Informed consent process**

The protocol and the informed consent were authorized by the La Sabana University Clinical Research Ethics Committee (CREC) which is ruled by the legal and ethical guidelines of the resolutions 008430 of 1993 and 002378 of 2008 of the Ministry of Social Protection (Colombia), World Conference on Harmonization for Good Clinical Practice of Institutions Conducting Investigation in Human Subjects and by the World Medical Assembly principles published in the Declaration of Helsinki, last review in 2008 [7-9].

A chat addressed to the volunteers was performed in order to explain the study in detail emphasizing in the type of medication to be used, planned dose, potential drug adverse reactions, blood volume to be collected at each study phase, the material to be used to collect the samples, the staff in charge of collecting them, diet restrictions to comply with, expected benefits for the community with the results and all the information they requested to freely decide their participation in the study. Subsequently, every one of them signed an informed consent form.

**Study design**

A randomized, open-label, two periods, two sequences, crossover design was used with an 8 days washout period between each period. Both products (Test and Reference) were administered 30 minutes after breakfast. The washout period was higher than 5 active ingredient half-lives. Three days before each period initiation, volunteers must refrain from medications, alcohol and any food or drink containing methylxantines. These restrictions prevailed during the whole sample collecting period. All volunteers were randomized to be allocated to the treatment sequence.

**Drug administration**

All volunteers received breakfast according to a regular diet in Colombia. Thirty minutes later, 2 tablets containing 50 mg of medication each (a total of 100 mg Desvenlafaxine succinate) were administered with 200 mL of water. During the entire experimental day, food intake completed 2600 calories divided into 85 g of protein, 78 g of fat and 390 g of carbohydrates. The plasma sample collecting team was confirmed by a physician and two licensed nurses. By means of a vacutainer⁴, a blood sample was collected by venipuncture in the superior limb immediately prior to administering the medication. Such sample was called zero time point sample. All volunteers received the Test product and the Reference product in each sequence. Thirteen blood venous samples were collected according to the following time points: 0; 2; 3; 4; 5; 6; 7; 8; 10; 12; 24; 36 and 48 hours. Samples were labeled for identification and centrifuged at 3000 rpm for 30 minutes. Plasma was transferred to a previously labeled tube and frozen at −20°C for later analysis.

**Analytical methodology**

Desvenlafaxine was quantified in plasma using a validated method. HPLC-UV method consisted of Desvenlafaxine extraction from 1.0 mL test plasma sample in a sedimentation centrifuge screw tap tube containing 60 µL of concentrated ammonium hydroxide. It underwent 20 seconds vortex stirring. 4 mL diethyl ether was added to this mixture and it underwent extra 60 seconds vortex stirring. Subsequently, it was centrifuged at 5000 r.p.m for 5 minutes. The organic layer was extracted in a glass tube and evaporated to dryness at 50°C under a gaseous nitrogen stream. Finally, the residue was reconstituted with 1.0 mL diluent solution and underwent 30 seconds vortex stirring. 100 µL were filtered and injected to the chromatographic system for RP-HPLC and UV detection analysis.

**Pharmacokinetic analysis**

The pharmacokinetic analysis was performed using WinNonlin 5.3 (Pharsight Corporation, Cary USA) software, by means of a non-compartmental analysis. Peak concentration (\(C_{\text{max}}\)) and the time to peak concentration (\(T_{\text{max}}\)) were obtained directly from the results of plasma concentrations, just as recommended by the FDA and the European Medicines Agency (EMA) for drug assessment [10-12].

Area under the curve, \(AUC_{\text{0-t}}\), between zero time point and the last time point with detectable concentrations was calculated through the trapezoidal rule, guaranteeing the calculation of at least 80% of the AUC with the last sample. The \(AUC_{\text{0-t}}\) was calculated as the C/Ke coefficient, being C the last detectable concentration and Kₑ the elimination constant, estimated as the slope obtained by linear regression from the points corresponding to the drug elimination phase through a linear regression of natural logarithm of concentrations [13]. Bioavailability-adjusted elimination constant (Kₑ), half-life (t½), clearance (Cl) and half mean residence time (MRT) were calculated after performing the non-compartmental analysis. The results of pharmacokinetic variables are summarized in Tables 3 and 4 with \(AUC_{\text{0-t}}\), \(AUC_{\text{0-}\infty}\), \(C_{\text{max}}\) and \(T_{\text{max}}\) values and the elimination time \(t_{\text{el}}\) of each one of the volunteers for each studied formulation.

**Statistical analysis**

An analysis of variance (ANOVA) was used to determine possible effects for each variation factor by sequence, period or subject. For this purpose, an F-test with a statistical significance level of 5% (\(p<0.05\)) was used. The statistical comparison of transformed pharmacokinetic parameters of both formulations was performed using the statistical software WinNonlin version 5.3. Pharmacokinetic parameters \(C_{\text{max}}\) and \(AUC_{\text{0-t}}\) were logarithmically transformed to conclude Bioequivalence if they met the 90% confidence interval within the 80-125% accepted limit range.

**Adverse events report**

Adverse events were recorded according to INVIMA guidelines Provision No. 1067/08, which defines them as serious or not serious and then, according to its definition, as likely, potential or not related relationship with the study medication. Since the sample size is not large, cases are informed as received from the investigation unit only and without any statistical estimation.

**Results**

Analytical results of active ingredient content, dose uniformity and dissolution test met the required specifications set by the Pharmaceutical Equivalence statement. Additionally, the comparative
dissolution profiles study showed a difference factor (f’) of 4 and a similitude factor (f2) of 77 Table 1.

The analytical method allowed the selective determination of Desvenlafaxine within the 5.0 to 270.0 ng/mL range of only 1 mL plasma. Method selectivity was proved through the analysis of six blank samples in which no interferences were detected between the analyte and the matrix endogenous compounds. Three Desvenlafaxine calibration curves were performed in human serum which showed to be linear within the 5 mg/mL to 300 ng/mL concentration range with linear correlation coefficients of 0.9977, 0.9986 and 0.9980; three Desvenlafaxine calibration curves were also performed in diluent solution, which showed to be linear in the same concentration range, with linear correlation coefficients of 0.9999, 0.9999 and 0.9998. Detection limit for the study was 1.7 ng/mL and quantitation limit was 5.0 ng/mL, which allows to perform a calibration curve in a wider concentration range. Precision expressed in terms of intratrat coefficient of variation was 14.8% for 15 mg/mL, 1.2% for 150 mg/mL and 3.0% for 270 mg/mL and interday coefficient of variation was 2.3% for 15 mg/mL, 5.0% for 150 mg/mL and 4.1 mg/mL for 270 mg/mL. All these values are below 15% for low, medium and high concentrations. The method is accurate as well, since the relative intratrat and intertest error values were also below 15% with -14.1%, 3.4% and 5.0% for 15 mg/mL, 150 mg/mL and 280 ng/mL, respectively and -5.2%, 6.4% and 9.1% for the same concentrations. Regarding the Desvenlafaxine recovery percent in human serum, percentages of 81.4% for 15 mg/mL, 84.7% for 150 mg/mL and 81.8% for 270 ng/mL were obtained with coefficients of variation of 5.5%, 6.0% and 8.4% respectively. Samples showed stability when stored at room temperature and refrigeration for 24 hours only, since after two storage days, evaluation showed evidence of degradation in 15 ng/mL concentration samples. Samples are stable in the freezing-thaw process for at least 3 cycles when stored at -20°C. It was therefore concluded that the quantitation method for Desvenlafaxine is selective, precise, accurate and shows good stability in human serum under analytical and instrumental laboratory conditions.

The study involved the participation of 24 healthy Colombian volunteers of both genders (50% women and 50% men) who completed both periods and were included in the pharmacokinetic and statistical analysis. Both treatments were well tolerated with the occurrence of adverse events considered as mild. The most commonly reported AE were nausea and pyrosis in 25% of the volunteers for both Test and Reference products (Table 2). Table 3 shows pharmacokinetic parameters obtained from all volunteers (mean ± SD) and Figure 1 shows comparative Bioavailability curves. The statistical analyses performed to determine Bioequivalence between the Test product Styma® produced by Tecnoquimicas S.A. and Pristiq® produced by Wyeth are shown in Table 4.

### Discussion

The reduction in costs of mental disorders treatment using multisource products showing the same quality and safety is a desired aim by governments interested in reducing the original drugs high costs. For this purpose, Bioequivalence studies assure the same pharmacokinetics for both generic and innovative products with no need of repeating clinical trials in new patients [14,15].

A rapid active ingredient release and a sudden increase of plasma concentration of Desvenlafaxine was discarded when performing the in vivo test with food presence according to WHO recommendations for products with a modified release system [16]. These findings are consistent with a study for assessment of Desvenlafaxine succinate pharmacokinetic changes when administered with or without food and that did not report significant changes in the molecule Bioavailability when interacting with food in the intestine [4]. The obtained values for Cmax and AUC0-t, either for the Test and Reference products are similar to those obtained in other studies [17].

This two periods, two sequences, crossover, single-dose design with healthy volunteers minimized the variability that is not due to formulation effects. The analytical method used was selective, precise, accurate and sensitive. The results indicate that the quantitation method for Desvenlafaxine is selective, precise, accurate and shows good stability in human serum under analytical and instrumental laboratory conditions.

Table 1: Test and Reference product physical-chemical tests results.

<table>
<thead>
<tr>
<th>Physical-chemical characteristic</th>
<th>Test Product</th>
<th>Reference Product</th>
</tr>
</thead>
<tbody>
<tr>
<td>Active ingredient</td>
<td>Test Product</td>
<td>Reference Product</td>
</tr>
<tr>
<td>assessment 90–110% of the labeled amount</td>
<td>50.1 mg</td>
<td>50.1 mg</td>
</tr>
<tr>
<td>Dosage Uniformity L1&lt;15</td>
<td>L1=2</td>
<td>L1=2</td>
</tr>
<tr>
<td>Dissolution test*</td>
<td>2 h&gt;30%</td>
<td>8 h</td>
</tr>
<tr>
<td>27.6%</td>
<td>45%</td>
<td>62%</td>
</tr>
</tbody>
</table>

* Dissolution time mean for 12 tablets

Table 2: Demographics of volunteers included in the pharmacokinetic and statistical analysis.

<table>
<thead>
<tr>
<th>Parameter (unit)</th>
<th>Test Product</th>
<th>Reference Product</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cmax (ng/mL)</td>
<td>215 ± 69.9</td>
<td>196.9 ± 58.2</td>
</tr>
<tr>
<td>AUC0-t (ng.h/mL)</td>
<td>3849.6 ± 1664.3</td>
<td>3605.4 ± 1412.1</td>
</tr>
<tr>
<td>AUC0-24h (ng.h/mL)</td>
<td>4233.7 ± 2076.8</td>
<td>3905.8 ± 1490.9</td>
</tr>
<tr>
<td>T1/2 (h)</td>
<td>6.3 ± 1.4</td>
<td>6.4 ± 1.7</td>
</tr>
<tr>
<td>K1 (1/h)</td>
<td>0.079 ± SD</td>
<td>0.068 ± SD</td>
</tr>
<tr>
<td>T1 (h)</td>
<td>10.5 ± SD</td>
<td>11.5 ± SD</td>
</tr>
</tbody>
</table>

Table 3: Pharmacokinetic parameters of desvenlafaxine succinate for Test product (Styma®) and Reference product (Pristiq®) followed by a 100 mg single oral postprandial dose. Mean and Standard Deviation are presented.

![Graphic representation of plasma concentration Vs Time Desvenlafaxine](image-url)
accurate and robust. All volunteers completed the study and only mild adverse events occurred when administering both formulations. The washout period was higher than the recommended 5 elimination half-lives and guaranteed the absence of carryover effect between periods. The Pharmaceutical Equivalence Statement and the similitude between dissolution profiles while estimating a non-difference factor ($f_1$)<15 and a similitude factor ($f_2$)>50 allowed to rate the in vitro quality attributes for both formulations. It was shown that mean values for AUC$_0-t$ and C$_{max}$ parameters were not significantly different between products and that the 90% confidence intervals for C$_{max}$ and AUC$_0-t$ for Test/Reference proportions remained within the ranges established by the regulatory entities for the Bioequivalence statement. The study sampling time was 48 hours, obtaining an AUC$_0-t$ greater than 80%, as suggested by the FDA published guideline [11].

Nichols et al [4] reported a Desvenlafaxine T$_{max}$ of approximately 6 hours in fasting conditions and approximately 2 to 4 additional hours when administered with food without affecting the apparent elimination half-life of oral dose except for the peak mean plasma concentration reached in postprandial conditions with a high fat content [4]. The lowest increase of peak plasma concentration in high fat content conditions was not clinically relevant [4]. In our study, there was no effect in the mean T$_{max}$ extension for both products following 6 hours when administered with food without affecting the apparent elimination half-life of oral dose except for the peak mean plasma concentration reached in postprandial conditions.

<table>
<thead>
<tr>
<th>Units</th>
<th>Ratio% T/R</th>
<th>Standard CI 90% (Test/Reference)</th>
</tr>
</thead>
<tbody>
<tr>
<td>ln C$_{max}$</td>
<td>108.49</td>
<td>103.58</td>
</tr>
<tr>
<td>ln AUC$_0-t$</td>
<td>104.46</td>
<td>97.96</td>
</tr>
<tr>
<td>ln AUC$_{0-∞}$</td>
<td>104.57</td>
<td>98.32</td>
</tr>
<tr>
<td>ln T$_{max}$</td>
<td>98.6</td>
<td>89.86</td>
</tr>
</tbody>
</table>

Table 4: 90% CI of logarithmically transformed pharmacokinetic parameters of two formulations containing Desvenlafaxine succinate (Test and Reference products) after its administration to healthy volunteers.

Conclusions

The Desvenlafaxine ER formulation Styma® manufactured by Tecnoquimicas S.A. (Test product) and Pristiq® manufactured by Wyeth (Reference product) have the pharmacokinetic parameters leading to the Bioequivalence statement between both formulations.

Acknowledgement

The test product is manufactured by Tecnoquimicas S.A. in Jamundí, Colombia under the trademarks Styma® and Desvenlafaxina MK® in Colombia, Desvenlafaxina MK® in Ecuador and Desvenlafaxteg TG® in Central America.

References