A New HPLC Method for Determination of Losartan in Human Plasma and its Application in Bioequivalence Studies

F. SHOKRANEH¹, A. DABIRSIAGHI¹* and N. ADIB²

¹Department of Pharmaceutics, Pharmaceutical Sciences Branch, Islamic Azad University (IAU), Tehran, Iran.
²Food and Drug Lab Research Center, Ministry of Health, Tehran (Iran).
*Address For Correspondence: Islamic Azad University (IAU), DrShariati St. Gholhak, Yakchel St. Tehran, Iran. 19395-6466
Email: alireza_dabirsiaghi@yahoo.com

(Received: February 04, 2012; Accepted: March 11, 2012)

ABSTRACT

A reliable, simple and sensitive reversed-phase high-performance liquid chromatographic method was developed for determination of losartan in plasma. Separation is achieved by HPLC after direct injection on a CN (250*4.6 mm) analytical column with a mobile phase composed of sodium hydrogen phosphate buffer - acetonitrile - tetrahydrofuran - methanol and phosphoric acid (0.1:5:4:21:69.9) V/V% adjusted to pH=9.9. Detection is by ultraviolet absorbance at 254nm. The flow rate was set at 0.6 ml/min. The lower limit of quantitation was 5 ng/ml. The intra and inter-day precisions (CV %) of the quality control samples were 0.57-5.31% and 0.21-4.52% respectively. The recovery of method was 92.25±2.19. The method was applied to a bioequivalence study in human.

Key words: Losartan, Human plasma, HPLC, Bioequivalence

INTRODUCTION

The renin-angiotensin (RAS) is a major regulator of blood pressure (BP). Losartan (COZAAR) is the prototype of a new class of orally active, non-angiotensin II receptor antagonists (ARBs) able to inhibit the renin-angiotensin system specifically and selectively without the agonistic effects of the peptide receptor antagonists. Approximately 14% of an oral dose of losartan is converted to the 5-carboxylic acid metabolite EXP 3174, which is more potent than losartan as an AT1-receptor antagonist. The metabolism of losartan to EXP 3174 and to inactive metabolites is mediated by CYP2C9 and CYP3A4. Peak plasma levels of losartan and EXP 3174 occur 1–3 hours after oral administration, respectively, and the plasma half-lives are 2.5 and 9 hours, respectively. The plasma clearances of losartan and EXP 3174 (600 and 50 mL/min, respectively) are due to renal clearance (75 and 25 mL/min, respectively) and hepatic clearance (metabolism and biliary excretion). The plasma clearance of losartan and EXP 3174 is affected by hepatic but
not renal insufficiency. Losartan should be administered orally once or twice daily for a total daily dose of 25–100 mg.3 Last studies showed that several HPLC, LC/MS/MS methods were used for determination of losartan and its metabolite in human plasma.4-12 The aim of this study was to develop a simple, rapid sensitive and reliable HPLC method with Ultraviolet detection for quantization of losartan in human plasma samples and to compare the bioavailability of two losartan tablets (50 mg) formulations (losartan from Iranian company, as a test formulation and COZAR, MSD, The Netherlands as a reference formulation). The method was validated according to procedures and acceptance criteria based on FDA guideline and recommendations of ICH, to provide enough selectivity, sensitivity and reliability in pharmacokinetic and bioequivalence studies.

MATERIALS AND METHODS

Hydrochloric acid, methanol (HPLC grade), ammonia, phosphoric acid, sodium hydrogen phosphate, acetonitrile, tetrahydrofurane, were purchased from Merc. Losartan and Thioridazine were USP reference standard.

Sample and standard solutions preparation

Blood samples were collected from individuals and centrifuged in order to separate the plasma. (Separated plasma was stored at -20 °C.). To a 0.5 ml aliquot of plasma, 1ml of internal standard (thioridazine) was added. 500 µl of 0.2 M HCL was used to acidify the solution and then vortexed and transferred in to the cartridge (Bond Elut™) CN which was prepared by using pure methanol and water in a proportion of 2:1. Initially 500µl distilled water and 10% methanol followed with 50µl of pure methanol was put through the cartridge. The internal standard and drug was then collected. The pH of the solution was made alkaline (pH =8) using ammonia/methanol in a proportion of 1:99 respectively and then dried under Nitrogen. The mobile phase was then poured in to the product in order to reduce it (The product was then reduced using the mobile phase (100µl)). 80µl of this solution was injected in to the HPLC with UV detector with the flow rate of 0.6ml/min and the column of CN (250*4.6mm). Stock solution of losartan (25 to 5000ng/ml) was prepared and diluted by blank plasma to obtain the final concentration (2.5 to 500ng/ml) of Losartan curve. Data from HPLC in defined circumstances was analyzed and evaluated.

Instrument and chromatographic conditions

Analyses were performed on younglin model ACME-900 pump equipped with Ultraviolet detector at wavelength of 254nm. Chromatography was performed at room temperature on a CN column (250*4.6mm). The mobile phase consisted of sodium hydrogen phosphate buffer - acetonitrile - tetrahydrofurane - methanol and phosphoric acid (0.1:5:4:21:69.9) V/V% adjusted to pH 9.9 using phosphoric acid. The flow rate was set at 0.6ml/min respectively.

Method validation

The method validation demonstrated the specificity, lower limit of quantification (LOQ), recovery, linearity, precision and accuracy of measurements.13 Specificity was investigated by analyzing six drug-free plasma samples for interference of endogenous compounds. For calibration curve five different concentrations of losartan (2.5 to 500ng/ml) in plasma were prepared by adding required volume of working solutions to blank plasma. Plasma calibration curve was prepared by taking area ratio of analyte to internal standard as Y-axis and concentration of analyte (ng/ml) as X-axis. Linearity of the standard curve was evaluated using least squares linear regression analysis. The limit of quantification (LOQ) was taken in this work at the lowest concentration standard affording accuracy and precision d”20%, using five plasma samples. The intra and inter-day precisions (CV %)of the assay procedure were determined by trice analysis of quality control plasma samples (50,200 and 500 ng/ml) at the someday and three different days. Recovery was determined by comparing the response of three pre-treated quality control plasma samples in three levels (25, 100 and 250 ng/ml) with the absolute peak area of un-extracted samples containing the same concentration of the drug as 100%.

Application

The validated method was used in bioequivalence study of losartan. It was an open,
one-centre, cross-over, randomized, three way and double-blind study to assess relative bioavailability of losartan in twenty-four healthy volunteers following single dose administration of losartan as 50 mg tablet (All subject gave informed consent to the work). The reference (COZAR, MSD, The Netherlands) and test (manufactured by Iranian company) product were used in this study. The blood collecting times were 0, 0.25, 0.5, 0.75, 1, 1.5, 2, 3, 4, 5, 6, 8, 12, 24, 36, 48 hour after oral administration of 50 mg losartan reference and test to fasting volunteers. The plasma samples were analyzed by the described method. The pharmacokinetic parameters like area under the plasma-concentration curve from zero to the last measurable losartan sample time and to infinity (AUC from and AUC infinite), maximum concentration ($C_{max}$), time to maximum concentration ($T_{max}$) were determined for the period of 0-48 hour.

RESULTS AND DISCUSSION

Some HPLC and LC-MS-MS methods have been developed for determination of losartan. The proposed method is suitable for losartan quantification in plasma samples. The based on a simple liquid – liquid extraction (LLE) followed by high – performance liquid chromatography with UV detector, using thioridazine as an internal standard, respectively. Blood samples were collected at specified time intervals, and the plasma separated and analyzed for losartan. Blank plasma and spiked plasma with losartan and internal standard are shown in Fig. 1. Retention time for the losartan and internal standard were 3.5 min and 8 min. The chromatograms also confirmed the complete separation. The calibration curve could be described by the equation:

$$\text{Conc} = 77\text{area}\% + (-6.1), \quad r^2 = 0.9921$$

The assay exhibited a linear dynamic range of 2.5 – 500 ng/ml, a run time of 20 min for each sample made it possible to analyze more than 70 samples per day. The limit of quantitation (LOQ) was 5 ng/ml. Intra and inter-day precisions (CV%)

![Fig.1. Chromatograms of (1) Blank human plasma; (2) Plasma spiked with thioridazine (rt=3.5min) and losartan (rt=8.0 min); (3) Human plasma after administration of losartan tablet](image-url)
The plasma samples from 24 health human volunteers were assayed with the validated method described above. The mean concentration – time curve is shown in figure 2.

Maximum plasma concentration (C_max) ranged from 104.53 to 176.44 ng/ml for test product and 121.03 to 192.60 ng/ml for reference product at 1.5 to 4 hour (T_max). Also the mean value of area under the concentration time curve (AUC₀-t) obtained was 1485.60 ng h/ml, AUC₀-∞ was found to be 2397.45 ng h/ml. No statistical differences were observed for C_max, T_max, AUC₀-∞ and other pharmacokinetic parametersand the test formulation is bioequivalent to the reference formulation for losartan.

The paper describes a rapid and reproducible HPLC method which enables the determination of losartan in plasma.

The main advantage of this method is the use of precipitation for purification, which is easily and fast in comparison with other purification and extraction methods. This HPLC method is reliable, reproducible and sensitive with respect to validation parameters. It can be used as an assay method in the study of losartan pharmacokinetics as well as bioavailability/ bioequivalence studies.
REFERENCES


