Manipulation of different ratio dependent approaches for determination of pravastatin in the presence of pioglitazone in pharmaceutical preparation

Abstract

Objective: This study aimed to develop three simple UV spectrophotometric methods for determination of pravastatin sodium in the presence of pioglitazone hydrochloride without previous separation.

Methods: Ratio spectra were developed for the interested methods. The first method is the ratio subtraction using 16 µg/mL of pioglitazone hydrochloride as a divisor. The second method is an amplitude modulation using the normalized spectrum of pioglitazone hydrochloride as a divisor (1 µg/mL) the peak amplitudes of ratio spectra were measured at isosbestic point 251.4 nm. The third method is the ratio difference using 14 µg/mL of pioglitazone hydrochloride as a divisor, where the peak amplitudes of ratio spectra were measured at 240 and 246.4 nm.

Results and discussion: The calibration curve is linear over the concentration range of 2-20 µg/mL, the proposed methods were validated according to International Conference on Harmonization (ICH) guidelines and successfully applied for the determination of pravastatin sodium in the presence of pioglitazone hydrochloride in their combined pharmaceutical formulation, Pravazon® capsules with high sensitivity.

Conclusion: The proposed three methods are simple, rapid, economical, accurate and precise for determination of pravastatin sodium in the presence of pioglitazone hydrochloride without previous separation.

Keywords: pravastatin sodium, pioglitazone hydrochloride, ratio subtraction, amplitude modulation, ratio difference

Introduction

Pravastatin sodium (PRV), (Figure 1A), is sodium (3R,5R)-7-[(1S,2S,6S,8S,8aR)-1,2,6,7,8,8a-hexahydro-6-hydroxy-2-methyl-8-[(S)-2-methylbutyryloxy]-1-naphthyl]-3,5-dihydroxyheptanoate. It is a selective and competitive inhibitor of 3-hydroxy-3-methylglutaryl-coenzyme A (HMG-CoA) reductase, the rate-limiting enzyme that converts HMG-CoA to mevalonate, a precursor of cholesterol. PRV is a member of the class of statins, used to treat hypercholesterolemia and related conditions and to prevent cardiovascular disease. It increases the number of hepatic low density lipoprotein (LDL) receptors on the cell surface to enhance uptake and catabolism of LDL. Secondly, PRV inhibits hepatic synthesis of very low density lipoprotein (VLDL), which reduces the total number of VLDL and LDL particles.

Pioglitazone hydrochloride (PGZ), (Figure 1B), is 5-[[4-(5-Ethyl-2-pyridinyl) ethoxy]phenyl]methyl]-2,4-thiazolidinedione hydrochloride. It is a thiazolidine-dione oral antidiabetic agent that improves insulin sensitivity. It is used in the treatment of type 2 diabetes mellitus. It is given orally as monotherapy, particularly in patients who are overweight and for whom metformin is contraindicated or not tolerated. Reviewing the literature on the new combination comprises PRV and PGZ in commercial dosage form, reveals only two reported methods for simultaneous determination of both drugs. The aim of this work is to develop simple, economic, rapid, sensitive, accurate and precise UV spectrophotometric methods for determination of PRV in the presence of PGZ without previous separation steps. These methods include; Ratio subtraction; Amplitude modulation; Ratio difference. Although this work is not advanced methods, but its value arises from the lack of any published method for determination of PRV in the presence of PGZ in their new combination.

Figure 1 Chemical structure of (A) pravastatin and (B) pioglitazone.
Instrument

Shimadzu UV/visible double beam 1800 Spectrophotometer (Tokyo, Japan) with 1 cm matched quartz cells and UV Probe software (2.35 ver.).

Materials and methods

Materials

Pure materials of PRV and PGZ were obtained as a gift sample from Hi Pharm Company, El-Obour City, Egypt. Their purity was found to be 99.52 and 99.61% for PRV and PGZ, respectively, according to the reported methods. Pravazon® capsules containing 10 mg of each drug per capsule (B.No. APB180816), manufactured by Accent Pharma, India, marketed by UPHA Pharmaceutical Mfg, India. Methanol, analytical grade (El-Nasr Company, Egypt).

Preparation of standard solutions

Standard solutions (100µg/mL) of PRV and PGZ were prepared by transferring accurately weighed 10mg of both drugs into two separate 100mL volumetric flasks, then dissolved in methanol and diluted to the mark with the same solvent.

Construction of calibration curves

Accurately measured aliquots equivalent to (0.20-2.00mL) of each drug were transferred from their standard solutions (100µg/mL) into two separate series of 10mL volumetric flasks and the volume of each flask was diluted up to the mark with methanol, to reach the concentration range of (2-20µg/mL). The absorption spectra of these solutions were measured in the range of 200 to 350nm against methanol as a blank.

Laboratory prepared mixtures

Accurate aliquots equivalent to (20-18µg) of PRV into a series of 1 ml volumetric flasks from its standard solution (100µg/mL) and portion equivalent to (180-20µg) of PGZ from its standard solution (100µg/mL) were added to the same flasks and volumes were completed to mark with pure methanol and mixed well.

Ratio subtraction method

The absorption spectra of the mixture were divided by the spectrum of 16µg/mL of PGZ (divisor) to give ratio spectra, then the constant was determined from the plateau region, then subtracted from the ratio spectra, the obtained spectra have the absorbance at isosorbptive point equal to the corresponding concentration of PRV.

Ratio difference method

The absorption spectra of the mixture were divided by the spectrum of 14µg/mL of PGZ. The peak amplitude of the ratio spectra was measured at 240 and 246.4nm. The Calibration graphs relating the difference in the two selected wavelengths to the corresponding concentrations of PRV were constructed, and the corresponding regression equation was computed.

Application to pharmaceutical formulation

The content of ten capsules was accurately weighed and mixed well. The amount of the powder equivalent to 10mg of PRV and 10mg of PGZ were weighed, dissolved in methanol by shaking for about 30min. The solution was filtered and transferred quantitatively into 100mL volumetric flask, the filtration system was evaluated to ensure that the filter does not adsorb any of the drugs. The volume was then diluted to the mark with methanol. Necessary dilutions were made to reach concentrations in the linearity range. The same procedures under the corresponding linearity were applied and the concentrations of PRV and PGZ were calculated from the corresponding regression equations.

Results and discussion

Simple spectrophotometric methods were developed for the determination of PRV in the presence of PGZ without previous separation. The absorption spectra of the resulting solution of PRV and PGZ which containing (2-20µg/mL) were measured and stored in the computer. Where, these absorption spectra of both drugs show severe overlapping and PGZ spectrum is extended more than PRV spectrum (Figure 2). The therapeutic importance of ROS and PRO in its combined pharmaceutical preparation was behind our interest in the development of simple methods for determination of PRV in presence of PGZ. The aim of the mixture is related to that PGZ exhibits a synergistic effect with statin drugs.
If the consultant determined directly from the spectra and subtracted, the new spectrum multiplied by $Y'$, the original spectrum of $X$ is obtained.

This can be summarized in the following equation:

$$\frac{X + Y}{Y'} = \left(\frac{X}{Y'}\right) + \left(\frac{Y}{Y'}\right)$$

(1)

By subtraction of the constant from equation (2):

$$\frac{X}{Y'} + \text{Constant} - \text{Constant} = \frac{X}{Y'}$$

(3)

By multiplication of equation (3) by a divisor ($Y'$):

$$(X/Y') \cdot Y' = X$$

(4)

The constant can be determined directly from the spectrum ($X+Y)/Y'$ by the straight line which is parallel to the wavelength axis in the region where $Y$ is extended.

The extended zero absorption spectrum of PGZ than zero absorption spectrum of PRV is necessary for application of this method as shown in (Figure 2). The absorption spectra of the mixture of PRV and PGZ were measured and recorded (Figure 3). The extension gives the constant value when we use a certain concentration of PGZ (16μg/mL) as a divisor (Figure 4), this constant can simply be determined directly from the obtained ratio spectrum in the region of PGZ absorption spectrum extension, and by subtraction of this constant the new ratio spectrum ($EZE$ spectrum / a divisor spectrum) was obtained. Good linearity is obtained in the concentration range of (2‒18μg/mL). The corresponding regression equation was computed.

Amplitude modulation method

The spectra of PRV and PGZ show the isoabsorptive point at zero spectrum and consequently retained as an isoabsorptive point at the ratio spectrum (Figure 5).

The absorbance of the zero order absorption spectrum for mixture of $X$ and $Y$ at isoabsorptive point as follow:

$$A_{mix} = \left[\frac{ax}{aY'}\right] + \left[\frac{aY}{CY}\right]$$

(1)

By dividing Eq. (1) with a normalized spectrum of Y as a divisor to get ratio spectrum with isoabsorptive point (At the same wavelength of the zero order) so the following equation was obtained:

$$A_{mix} = \left[\frac{ax}{aY'}\right] + \left[\frac{aY}{CY}\right]$$

(2)

$$A_{mix} = \left[\frac{ax}{aY'}\right] + \text{Constant}$$

(3)

Where, ($P_m$) is the amplitude of the ratio spectrum of the mixture, ($P_x$) is the amplitude of component $X$ and ($P_y$) is the amplitude of component $Y$. i.e. the recorded amplitude at the isoabsorptive point of the ratio spectrum is equal to the sum of amplitude corresponding to $X$ and that of $Y$.

The amplitude representing the component $Y$ ($P_y$) was the constant

$$\left[\frac{aY}{aY'}\right]$$

and it can be measured directly from the spectrum at the straight line that is parallel to the wavelength axis in the region where Y spectrum is extended.

Since, we use a normalized divisor of Y (1μg/mL)

$$P_y = \left[\frac{aY}{aY'}\right]$$

(4)

$$P_x = \left[\frac{ax}{aY'}\right]$$

(5)

$\therefore$ The recorded amplitude of the constant was modulated to concentration so it was representing the concentration of $Y$ [$CY$, (CRecorded of $Y$)].

For determination of amplitude of $X$ in the mixture, If we subtract the measured value of the constant from that of the mixture at isoabsorptive point of the ratio spectrum Eq. (2);

$$P_x = P_m - PY$$

(6)

$\therefore$ At that isoabsorptive point $a_x = a_y$ and normalized divisor of Y $CY = 1μg/mL$. 

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\[ P_x \left( \frac{ax}{aY} \right) \]

\[ P_x = \left[ \frac{C_x}{CY} \right] \]

This obtained amplitude of ratio spectrum was modulated to concentration and it was representing the concentration of X \([C_x]\), \(C_{\text{Recorded}}\) of X).

The corresponding concentration of X or Y could be calculated by using the following regression equation:

\[ C_{\text{Recorded}} = \text{Slope} \cdot C + \text{intercept} \]

Slope was found to be approximately one and intercept almost zero.

Where, \(C_{\text{Recorded}}\) represents the recorded amplitudes corresponding to the concentrations of either X or Y that obtained from the ratio spectrum using a normalized spectrum of Y (1μg/mL) as a divisor and C represents the corresponding concentration of X or Y.

This method based on the same principle as the ratio subtraction method with simple modification step [use the normalized spectrum (1μg/ml) as a divisor] (Figure 6).

Good linearity is obtained in the concentration range of (2–18μg/mL). The corresponding regression equation was computed.

Validation of the method

The proposed method was tested in the linearity range, limit of detection (LOD), limit of quantification (LOQ), accuracy and precision according to International Conference on Harmonization (ICH) guidelines.\(^{14}\)

Linearity and range

According to the described theory of ratio subtraction method, the linearity of PRV was determined at 237nm. According to the theory of amplitude modulation method, the linearity of PRV was determined by measuring the peak amplitudes of ratio spectra at the isoabsorptive point of 251.4nm.

According to the experimental conditions of the ratio difference method, the calibration graphs for the method was constructed by plotting the differences of peaks amplitude at 240 and 246.4nm against the concentration which related only to PRV. The calibration graphs for all proposed methods were rectilinear over the concentration range of 2-20μg/mL for PRV with high values of the correlation coefficient, the data is supplied in Table 1.

![Figure 5 Zero-order absorption spectra of PRV (10μg/ml), PGZ (10μg/ml) and mixture of both drugs (5μg/ml of each one)](image)

![Figure 6 Ratio spectra of laboratory prepared mixtures of PRV and PGZ using normalized spectrum (1μg/ml) of PGZ as a divisor](image)

### Table 1 Spectral and validation data for determination of PRV in presence of PGZ by the proposed methods

| Parameters          | Ratio subtraction | Amplitude modulation | Ratio difference |
|---------------------|-------------------|----------------------|------------------|
| Wavelength (nm)     | 237               | 251.4                | 240 and 246.4    |
| Linearity range (μg/mL) | 2—20          | 2—20                | 2—20            |
| LOD (μg/mL)         | 0.241             | 0.329               | 0.187            |
| LOQ (μg/mL)         | 0.729             | 0.988               | 0.567            |
| Regression equation |                   |                      |                  |
| Slope               | 0.0523            | 1.0044              | 0.0434           |
| Intercept           | 0.007             | -0.0178             | 0.0006           |
| Correlation coefficient (r) | 0.9998        | 0.9999              | 0.9998           |
| Accuracy (%R)*      | 99.63             | 100.21              | 100.33           |
| Precision (%RSD)*   |                   |                      |                  |
| - Repeatability (intra-day) | 0.971        | 0.675               | 0.752            |
| - Intermediate precision | (Inter day)   | 1.014               | 0.806            | 0.822          |

*Average of three determinations of three concentration levels.

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Limit of detection (LOD) and limit of quantification (LOQ)

LOD and LOQ were calculated according to ICH guidelines from the following equations:

\[
LOD = \frac{3.3 \cdot S_a}{\text{slope}}
\]

\[
LOQ = \frac{10 \cdot S_a}{\text{slope}}
\]

Where \( S_a \) is the residual standard deviation of a regression line.

LOD and LOQ values of PRV and PGZ for each method were listed in Table 1.

Selectivity

The selectivity of the methods was achieved by the analysis of different laboratory prepared mixtures of PRV and PGZ within the linearity range. Satisfactory results listed in Table 2, and the results of the standard addition technique Table 3 prove that the proposed methods can selectively analyze PRV without any interference from the other drug (PGZ) or the excipients.

Accuracy

Accuracy of the proposed methods, calculated as the mean percent recovery (\%R), was assessed by applying the proposed procedures for triplicate determination of three concentration levels covering the specified range for each drug (6, 10 and 16μg/mL). The concentrations were obtained from the corresponding regression equations and the mean percent recoveries, shown in Table 1, indicate the accuracy of the proposed methods. Accuracy of the methods was further assured by the use of the standard addition technique. It was performed by the addition of known amounts of pure PRV and PGZ to known concentrations of the pharmaceutical preparation and the resulting mixtures were assayed, and the results obtained were compared with the expected results (Table 3). The good recoveries of the pure added PRV suggested good accuracy of the proposed methods.

Precision

Precision of the proposed methods, calculated as percent relative standard deviation (\%RSD) of the percent recoveries, was checked by applying the proposed procedures for triplicate determination of three concentration levels covering the specified range for each drug (6, 10 and 16μg/mL) in the same day (intra-day analysis) for repeatability and on three different days (inter day analysis) for intermediate precision. The results in Table 1 indicate precision of the method.

Application to pharmaceutical formulation

The proposed methods were applied for the determination of PRV and PGZ in their combined pharmaceutical formulation, Pravazon® capsules. Satisfactory results were obtained in good agreement with the label claim, and the results of the standard addition technique indicate no interference from excipients and additives (Table 3).

Statistical analysis

Table 4 showed statistical comparison of the results obtained by the proposed methods and the reported method in their pure form. The calculated \( t \) and \( F \) values were less than the theoretical ones indicating that; there was no significant difference between the proposed and the reported methods with respect to accuracy and precision.

Table 2 Determination of PRV in presence of PGZ in the laboratory prepared mixtures by the proposed methods.

| Method          | Concentration taken (µg/mL) | Concentration found (µg/mL) | % Recovery |
|-----------------|----------------------------|-----------------------------|------------|
| Ratio Subtraction | 18 PRV, 18 PGZ            | 17.88 PRV                   | 99.32      |
|                 | 16 PRV, 16 PGZ            | 16.04 PRV                   | 100.26     |
|                 | 14 PRV, 14 PGZ            | 14.07 PRV                   | 100.52     |
|                 | 12 PRV, 12 PGZ            | 12.14 PRV                   | 101.18     |
|                 | 10 PRV, 10 PGZ            | 10.1 PRV                    | 100.96     |
|                 | 8 PRV, 8 PGZ              | 8.01 PRV                    | 100.14     |
|                 | 6 PRV, 6 PGZ              | 6.04 PRV                    | 100.7      |
| Mean±%RSD       |                           | 100.44±0.611                |            |
| Amplitude modulation | 18 PRV, 18 PGZ          | 18.06 PRV                   | 100.34     |
|                 | 16 PRV, 16 PGZ            | 16.15 PRV                   | 100.91     |
|                 | 14 PRV, 14 PGZ            | 13.97 PRV                   | 99.8       |
|                 | 12 PRV, 12 PGZ            | 12.07 PRV                   | 100.57     |
|                 | 10 PRV, 10 PGZ            | 10.0 PRV                    | 100        |
|                 | 8 PRV, 8 PGZ              | 8.03 PRV                    | 100.55     |
| Mean±%RSD       |                           | 100.31±0.399                |            |
| Ratio difference | 18 PRV, 18 PGZ            | 17.91 PRV                   | 99.5       |
|                 | 16 PRV, 16 PGZ            | 15.93 PRV                   | 99.56      |
Table 3 Determination of PRV and PGZ in Pravazon® capsules by the proposed methods and application of standard addition technique

| Proposed methods         | Pravazon® capsules | Standard addition technique |
|--------------------------|--------------------|-----------------------------|
|                          | % Recovery±%RSD     | Taken (µg/mL) | Pure added (µg/mL) | Pure found* (µg/mL) | % Recovery |
| Ratio Subtraction        | PRV                | PRV          | PGZ          | PRV          | PGZ          | PRV          | PRV          |
|                          | 99.44±1.706        | 4            | 4            | 4            | 100.75       |
|                          | 99.44±1.706        | 6            | 6            | 6.13         | 101.17       |
|                          | 99.44±1.706        | 10           | 10           | 10.06        | 100.6        |
| Mean±%RSD                | 99.44±1.706        | 4            | 4            | 4.06         | 101.5        |
| Amplitude modulation     | 99.48±0.858        | 4            | 4            | 6.03         | 100.5        |
|                          | 99.48±0.858        | 6            | 6            | 9.94         | 99.6         |
|                          | 99.48±0.858        | 10           | 10           | 9.92         | 99.2         |
| Mean±%RSD                | 99.48±0.858        | 4            | 4            | 3.97         | 99.25        |
|                          | 99.48±0.858        | 6            | 6            | 6.07         | 100.17       |
|                          | 99.48±0.858        | 10           | 10           | 9.92         | 99.2         |
| Ratio difference         | 100.12±1.371       | 4            | 4            | 9.92         | 99.2         |
|                          | 100.12±1.371       | 6            | 6            | 6.07         | 100.17       |
|                          | 100.12±1.371       | 10           | 10           | 9.92         | 99.2         |
| Mean±%RSD                | 100.12±1.371       | 4            | 4            | 3.97         | 99.25        |

* Average of three determinations

Table 4 Statistical comparison between the results obtained by the proposed methods and the reported methods for the determination of PRV* and PGZ* in pure powder form

| Parameters        | Ratio subtraction | Amplitude modulation | Ratio difference | Reported method |
|-------------------|-------------------|----------------------|------------------|-----------------|
|                   | PRV               | PRV                  | PRV              | PRV             |
| n                 | 5                 | 5                    | 5                | 5               |
| Mean %R           | 99.63             | 100.21               | 100.33           | 100.07          |
| %RSD              | 1.018             | 0.806                | 0.635            | 0.564           |
| Student's t-test  | (2.306)           | (3.068)              | (2.306)          | (3.068)         |
| (c)               | 1.125             | 0.907                | 0.689            | —               |
| F value           | (6.388)           | (6.388)              | (6.388)          | —               |
| (6.388)©          | 1.305             | 2.651                | 1.969            | —               |

Conclusion

Although these proposed methods are not advanced, their values arise from the lack of any published method for determination of PRV in the presence of PGZ in their new combination. The proposed methods are simple, rapid, economical, accurate and precise and can be used for determination of PRV in the presence of PGZ in pure form as well as in pharmaceutical dosage form without interference from excipient and no need for previous separation steps.

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Conflict of interest

The authors declare there is no conflict of interest.

References

1. Sweetman S. Martindale: the complete drug reference. 36th ed. London: The Pharmaceutical Press; 2009.
2. Moffat AC, Osselton MD, Widdop B. Clarke’s analysis of drugs and poisons. 3rd ed. London: The Pharmaceutical Press; 2005.
3. Lennernas H, Fager G. Pharmacodynamic and pharmacokinetics of the HMG–CoA reductase inhibitors. Clin Pharmacokinet. 1997;32(5):403–425.
4. Nissen S, Nicholls S, Sipahi I, et al. Effect of very high-intensity statin therapy on regression of coronary atherosclerosis: the Asteroid trial. JAMA. 2006;295(13):1556–1565.
5. The Merck index. 14th ed. USA: Whitehouse Station, Merk & Co Inc. NJ; 2006.
6. Olemy A. Simultaneous UV Spectrophotometric Determination of Pravastatin Sodium and Pioglitazone Hydrochloride in Pharmaceutical Preparations. J Adv Pharm Res. 2017;1(3):143–149.
7. Balaji S, Kattebona S. Development of spectrophotometric method for determination of pravastatin sodium in bulk and tablet formulations. Int J Pharm Tech Res. 2009;1(4):1017–1019.
8. Ali MY, Swamy PV, Bargaonkar P, Raju SA. UV–spectrophotometric determination of pioglitazone in pharmaceutical dosage forms. Int J Chem Sci. 2008;6(4):2062–2065.
9. Khalid AM, Mohamed WI, Ahmad AM, et al. Simple stability indicating ultraviolet spectroscopic methods for the estimation of ezetimibe in presence of its alkaline degradation product. ACAIJ. 2016;16:382–393.
10. Khalid AM, Mohamed WI, Ahmad AM, et al. A comparative study of different approaches for stability indicating determination of tizanidine in presence of its oxidative degradation product. Int J Pharm Pharm Res Human. 2016;5:166–195.
11. Tonker T, Yanan Wang, Willeke de Haan, et al. Pioglitazone Decreases Plasma Cholesteryl Ester Transfer Protein Mass, Associated With a Decrease in Hepatic Triglyceride Content, in Patients With Type 2 Diabetes. Diabetes Care. 2010;33(7):1625–1628.
12. Leonhardt W, Andreas Pfützner, Jürgen Müller, et al. Effects of pioglitazone and/or simvastatin on low density lipoprotein subfractions in non-diabetic patients with high cardiovascular risk: A sub-analysis from the PIOSTAT study. Atherosclerosis. 2008;201(1):155–162.
13. Wei J, Ma C, Zhang Y, et al. Synergistic effects of pravastatin and pioglitazone in renal tubular epithelial cells induced by transforming growth factor-β1. Cell Biology International. 2007;31(5):451–458.
14. International Conference on Harmonization. Validation of analytical procedure: text and methodology: Q2 (R1). International Conference on Harmonization, Geneva, Switzerland; 2005.
15. International Conference on Harmonization. Validation of analytical procedure: text and methodology: Q2 (R1). International Conference on Harmonization, Geneva, Switzerland; 2005.