INTRODUCTION

Glimepiride is a medium-to-long acting sulfonylurea anti-diabetic drug. Pioglitazone is a prescription drug of the class thiazolidinedione (TZD) with hypoglycemic (antihyperglycemic, antidiabetic) action. Glimepiride is official in US Pharmacopoeia, which describes the HPLC method for the determination of either in raw material or in pharmaceutical formulations. Pioglitazone is not official in any pharmacopoeia. The literature contains several methods for the determination of glimepiride in pharmaceutical dosage forms, including liquid chromatography and derivative spectroscopy. Glimepiride related substances as well as degradation pathway methods were reported. Similarly degradation behavior of pioglitazone method and stability indicating assay method was also available. Some assay method is reported in the combination tablets. No stability indicating HPLC method has been reported yet for the simultaneous determination of glimepiride and pioglitazone in combination drug product. By seeing the glimepiride and pioglitazone degradation behavior it is essential and worth full to develop a stability indicating HPLC method. This combination drug product is marketed by many of the pharmaceutical company but till now no method is available to determine the said drug in the routine quality control and stability sample analysis. It is mandatory to develop a stability indicating assay method for the unstable molecule like glimepiride and pioglitazone. Additionally to prove the selectivity of the method, glimepiride major degradation impurity of related compound B and related compound C was injected and estimated in the combination tablet. A single HPLC method is described in this paper for the simultaneous estimation of pioglitazone, glimepiride, glimepiride related compound B and related compound C.

EXPERIMENTAL

Materials and reagents

Pharmaceutical grade standards of pioglitazone (5-(4-[2-(5-ethylpyridin-2-yl)ethoxy]benzyl)thiazolidine-2,4-dione) and glimepiride (3-ethyl-4-methyl-N-(4-[N-(1r,4r)-4-methylcyclohexyl(carbamoyl)sulfamoyl]phenethyl)-2-oxo-2,5-dihydro-1H-pyrrrole-1-carboxamide) were kindly supplied by M/S Pharma Lab (Baddi, India). Glimepiride related compound B (3-ethyl-4-methyl-2-oxo-N-[2-[4-sulphamoyl(carbamoyl)sulfamoyl]ethyl]2,3-dihydro-1H-pyrrrole-1-carboxamide) and related compound C (Methyl [4-[2-[([3-Ethyl-4-methyl-2-oxo-2,3-dihydro-1H-pyrrrole-1-y]carbonyl)amino]ethyl[phenyl]- sulphonyl]carbamate) were purchased from LGC standards (India). Chemical structures are shown in Figure 1. Commercially available combination tablets containing 15mg of pioglitazone, 2mg of glimepiride and 500mg of metformin HCl (PRICHEK GMP®-manufactured by Indoco Rem) were purchased. HPLC grade acetonitrile, analytical reagent grade potassium dichromate and ortho phosphoric acid were obtained from Rankem (India). Millipore water was used prepared from Milli-Q plus water purification system.

Chromatographic conditions

The HPLC system was a Waters 2695 binary pump plus auto sampler and a 2996 photo diode array as well as 2487 UV detector.

The chromatographic column used for separation was a Zorbax cyano, 250 x 4.6mm with 5 microns particles. Mobile phase consisted solution- A and solution-B. Solution-A is 20 mmol L⁻¹ potassium dihydrogen phosphate, pH adjusted to 3.2 using dilute ortho phosphoric acid. Solution-B is acetonitrile.

The gradient program was as follows: (time (min)/% solution B: 0.01/20, 13/20, 15/80, 40/50, 55/80, 60/80, 63/20, 70/20. Mixture of acetonitrile and water (4:1, v/v) were used as sample diluent. The flow rate of the mobile phase was 0.8mL min⁻¹. The column temperature was maintained at ambient and the detection was monitored at a wavelength of 230nm. The desired LOQ value of impurities were achieved by using 25 µL of injection volume.

System suitability solution

Stock solutions of glimepiride related compound B, related compound C and glimepiride (1000µg mL⁻¹) were prepared by dissolving appropriate
amounts. System suitability solutions of 0.2 µg mL⁻¹ of related compound B and related compound C, 0.5 µg mL⁻¹ of glimepiride were prepared from above stock solutions.

Standard preparation

A standard solution containing 750 µg mL⁻¹ of pioglitazone and 100 µg mL⁻¹ of glimepiride were prepared by dissolving appropriate amount of pioglitazone and glimepiride standard in diluent.

Preparation of sample solution

Twenty tablets were weighed and powdered with the help of mortar and pestle. An amount of powder tablets equivalent to 10 mg glimepiride (75mg of pioglitazone) were transferred to a 100 mL volumetric flask, 60 mL of diluent were added and kept on a rotary shaker for 10 min to disperse the material completely, then sonicated for 10 min (during the sonication the bath temperature was maintained at 25°C) and diluted to 100 mL with diluent. The concentration of pioglitazone and glimepiride was 750 µg mL⁻¹ and 100µg mL⁻¹. The resulting solution was centrifuged at 10000 rpm for 5 min. The supernatant solution was used for estimation of pioglitazone, glimepiride and glimepiride impurities.

RESULTS AND DISCUSSIONS

Optimization of chromatographic method

The HPLC method is optimized with a view to develop a stability indicating nature. A stability indicating method should accurately measure the active ingredients, without interference from degradation products and sample matrixes. Since pioglitazone and glimepiride is having a degradation nature gradient method is preferred over isocratic to get a complete degradation product as well as good resolution between close eluting compounds. To develop a method the initial trials were taken with pure drug of pioglitazone and glimepiride spiked with glimepiride related compound B and glimepiride related compound C was used a sample solution. Different buffer pH (2-7), different solvent system containing methanol and acetonitrile was tried. The reverse phase column chemistry of C18 is applied for preliminary trial. The good separation was achieved with gradient program contain solution A (phosphate buffer at pH 3.2) and solution B (acetonitrile) with a flow rate of 0.8mL min⁻¹. To prove the stability indicating nature of the method all forced degradation samples were injected in the optimized conditions, but the peak purity of glimepiride and pioglitazone got failed due to some degradation compound elution at the same RT of the peak pioglitazone and glimepiride peak. To check this some gradient adjustment, column temperature and flow rate was tried, but these trials were not come up with the good result. To rectify the problem, different column was tried initially C8 column was selected, but know compound itself got merged together in this column chemistry. While using phenyl column one degradation peak has come out from the glimepiride peak and glimepiride peak purity was found satisfied but pioglitazone peak purity was remain same. Finally the cyano column was used for the development; the main base degradation peak was come out with the more than 2.0 resolutions from pioglitazone peak. For our knowledge this is the first method was reported with the these many degradation peak is possible to form in the said combination drug product and all the degradation peak as well as known component was got very good resolution. All pioglitazone, glimepiride, glimepiride related compound B and related compound C was found adequate response at 230nm. In case of stressed sample, chromatogram was extracted with the entire range of 200-400nm to check a new impurity at different wavelength but no extra peak was found except at 230nm wavelength observed peaks. The required LOQ value of glimepiride related compound B and related compound C was found by using 100 µg mL⁻¹ of glimepiride sample preparation with 25µL injection volume. During the development it was observed that the formation of related compound B is very fast and to get a consistent result the fresh sample preparation was prepared and used, the sonicator both temperature was maintained less than 25°C while preparing the sample solution. The critical close eluting impurity of glimepiride related compound B and related compound C was found better resolution than the current USP monograph glimepiride tablet method. System suitability chromatogram is shown in (Figure. 2). This is proving that in all the aspect the developed method is better than the till now reported methods.

![Figure 2](image)

**Figure 2.** System suitability chromatogram (contain glimepiride, glimepiride related compound B, glimepiride related compound C).

Method validation

The developed chromatographic method was validated for system suitability, selectivity, specificity, linearity, precision, accuracy, LOD, LOQ and robustness as per ICH guideline.¹⁵

System suitability

The observed retention times (RT) and relative retention times of the analytes are presented in Table 1. The resolution between close eluting pair of glimepiride related compound B and glimepiride related compound C was set as the system suitability parameter (> 6.0). Also the RSD of the peak area of pioglitazone and glimepiride was calculated.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Pioglitazone</th>
<th>Glimepiride</th>
<th>Glimepiride related compound B</th>
<th>Glimepiride related compound C</th>
</tr>
</thead>
<tbody>
<tr>
<td>% RSD</td>
<td>1.1</td>
<td>1.3</td>
<td>4.1</td>
<td>3.2</td>
</tr>
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<td>Retention time</td>
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<td>USP resolution</td>
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<td>-</td>
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<td>USP theoretical plates</td>
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<td>8012</td>
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</tr>
</tbody>
</table>

Specificity and selectivity

Specificity of the developed method was assessed by performing forced degradation studies.

The specificity of the developed HPLC method was determined in the presence of it’s degradation products and other sample matrix. Forced degradation studies were performed on tablet sample to provide an indication of the stability-indicating property and specificity of the proposed method. The sample solutions were subjected to acid and base hydrolysis (using 0.1N HCL and 0.1N NaOH, respectively for 2h), to oxidation (using 3% H₂O₂ for 2h) and to UV radiation (254nm for 48h).

When the drug was exposed to acid and peroxide condition minor degradation was observed but in case of base condition major degradation was observed. Related compound B was increased in all acid, base and peroxide stressed sample but related compound C was found only in peroxide condition. Chromatograms are shown in (Figure. 3). The said drugs are stable to the effect of photolysis, no degradation was observed. In all stressed samples peak purity were found within acceptable limit (purity angle is less than purity threshold) indicating the specificity of the method. Results are shown in (Table 2). To prove the selectivity of the method metformin hydrochloride was injected in the optimized method, no interference and overlap was found with the respect to pioglitazone, glimepiride and glimepiride impurities peaks. Chromatogram is shown in (Figure. 4).

![Figure 3](image)

**Figure 3.** Chromatograms of the stressed sample. A) Acetone (2h), B) Base (0.1N NaOH) (2h), C) Base (5% H₂O₂) (2h), D) UV (254nm) (2h), E) Photolysis (254nm) (2h).

![Figure 4](image)

**Figure 4.** Chromatogram of pioglitazone, glimepiride and glimepiride impurities peak in the optimized condition.
The authors would like to thank M/S Pharma Lab (Baddi, India) for providing the standards.

**ACKNOWLEDGEMENTS**

A single reversed phase stability indicating RP-HPLC method has been established for simultaneous estimation of pioglitazone, glimepiride and glimepiride related compound B and related compound C from combination drug product. The method was fully validated and the data found to be satisfactory for all the method validation parameters tested. The developed method can be conveniently used by a quality control department to determine the said compound in routine analysis and commercial sample purity check.

**CONCLUSIONS**
REFERENCES