IN VITRO COMPARISON OF DIFFERENT BRANDS OF CETIRIZINE 2HCl TABLETS BY SPECTROSCOPY AND THIN LAYER CHROMATOGRAPHY (TLC) IN LOCAL MARKET OF LAHORE, PAKISTAN

SAJID MAHMOOD¹, AMMARA RAFIQ, ZAHEER AHMED, MUHAMMAD AJMAL

ABSTRACT. The objective of the present study is to investigate and compare the efficacy, physiochemical equivalence and purity of API's of different brands of cetirizine 2HCl tablets (10mg) with that of multinational brands available in local market of Lahore, Pakistan. The present work also provides awareness to allergy patients about most effective local brands with reasonable prices, without any health risk. Spectrophotometric method is used for chemical assay, TLC technique is subjected for the purity of API's and physiochemical parameters is employed i.e. weight variation, hardness, friability, thickness, disintegration time and dissolution study as well as price fluctuation in PK rupees. The results of all physiochemical parameters and chemical assay are found to be within limit and meet to the pharmacopeial standards. TLC technique is developed to check the quality and purity of API, all the active ingredients showed the similar Rf-values without any impurity (0.38-0.39). From the test results and plots, it is observed and concluded that an economical and quality products can be prescribed for allergy patients whether they are manufactured by the local or multinational pharmaceutical companies without any health risk.

Index Terms: Cetirizine 2HCl; comparative analysis; spectroscopy; TLC; physiochemical parameters.

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1. Introduction

Tablets and capsules represent unit dosage forms whereas liquid oral dosage forms such as syrups, suspensions, emulsions, solutions and elixirs usually contain one dose of medication in 5 to 30mL. Such doses are erratic by a factor ranging from 20 to 50% when the drug is self-administered by the patient [1]. The oral route of drug administration is the most important method for systemic effects. Solid oral dosage forms (tablet and capsule) are the preferred class of products of the two forms, the tablet has a number of advantages such as the tablet is an essentially tamper proof dosage form [2, 3]. Cetirizine is the carboxylated metabolite of hydroxyzine, and has high specific affinity for histamine H-receptors. Cetirizine dihydrochloride (CZ) is (RS)-2-[(4-chlorophenyl) phenyl methyl] piperazine-1-yl]ethoxy] acetic acid dihydrochloride. It is used for the symptomatic relief of allergic conditions including rhinitis and chronic urticarial [4, 5].

Literature survey shows comparative study of different brands of the same generic of pharmaceutical dosage form, and have indicated that pharmaceutical evaluation of different brands are as important as biological and clinical equivalency. It is generally evaluated that many drugs that are manufactured in developing countries are implicated to be substandard [6]. To improve the quality of health products and to minimize the health risk factors. It is necessary to monitor all the pharmaceutical services in a regular basis that promoting the conditions and providing information on the base of which the people become enable to make healthy choices and they can make correct decisions about their health. Many tests are frequently applied to tablet dosage forms to render their optimum therapeutic effects. The technique of optimization is well reported in the literature for the development of tablet formulations [7, 8, 9]. The purpose of carrying out optimization is to select the best possible formulation from a pharmaceutical as well as consumer point of view.

In the context above we have made an attempt to compare all the physical and chemical parameters including purity of API’s of five different brands of cetirizine 2HCl tablet available in local market of Lahore, Pakistan with that of multinational brand on the basis of GMP, GLP and Pharmacopial standards. Since no work has so far been carried out in the said perspective. Therefore we choosed this topic and to decide to provide awareness to allergy patients about most effective local brands with reasonable prices, without any health risk.

2. Materials and Methods

Reference cetirizine 2HCl was a kind gift sample from Jawa pharmaceuticals (pvt) Ltd. Lahore, Pakistan. Different brands of cetirizine 2HCl were obtained from different retail pharmacies of Lahore (Pakistan) and their qualitative and quantitative tests are carried out.

Zyrtec (UCB Farchim, S.A, Switzerland)
Alergo (Pharmix laboratories)
In vitro comparison of different brands

Cetrix (Saffron Pharmaceuticals)
Instazin (Friends Pharma)
Tryzin (Pulse pharmaceuticals).

3. Experimental

Spectrophotometer (Shimadzu UV-1800), Disintegration apparatus (IPL - 27 model) Hot plate with magnetic stirrer (78-1) Hardness tester (PAK/HT-0607), Friability apparatus (Betatek EF-2), Weighing balance (Japan), micropipette, Hydrochloric acid (Merck Germany), and freshly prepared distilled water are used for this study. Reasonable availability of medicine requires that a medicine should fulfill the clinical needs of a patient for an adequate period of time, at the lowest cost to them and their community. Various pharmaceutical parameters were employed as in USP 31 (2008), i.e. thickness, hardness, weight variation, friability, disintegration, and dissolution, to test the different brands of cetirizine hydrochloride tablets available.

3.1. Price Fluctuation. The most effective local brands with reasonable prices, without any health risk also provide awareness to the allergic patient in our community. Label information and price fluctuation among different brands of cetirizine 2HCl are given in Table 1.

<table>
<thead>
<tr>
<th>Sr No</th>
<th>Product Name</th>
<th>Manufactured by</th>
<th>Batch No</th>
<th>Mfg date</th>
<th>Exp date</th>
<th>Price/10 tab</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Zyrtec</td>
<td>UCB Farchim, S.A, Switzerland</td>
<td>HZXAP</td>
<td>04-2018</td>
<td>04-2020</td>
<td>42.00</td>
</tr>
<tr>
<td>2</td>
<td>Alergo</td>
<td>Pharmix laboratories (Pvt) Ltd</td>
<td>C824</td>
<td>03-2018</td>
<td>02-2021</td>
<td>39.00</td>
</tr>
<tr>
<td>3</td>
<td>Cetrix</td>
<td>Saffron Pharmaceuticals (Pvt) Ltd</td>
<td>106</td>
<td>11-16</td>
<td>10-19</td>
<td>28.00</td>
</tr>
<tr>
<td>4</td>
<td>Instazin</td>
<td>Friends Pharma (Pvt)Ltd</td>
<td>18IN021</td>
<td>08-2018</td>
<td>08-2021</td>
<td>42.00</td>
</tr>
<tr>
<td>5</td>
<td>Tryzin</td>
<td>Pulse pharmaceuticals (Pvt)Ltd</td>
<td>1298</td>
<td>11-2018</td>
<td>10-2021</td>
<td>30.00</td>
</tr>
</tbody>
</table>

3.2. Thickness and Diameter. Thickness of tablet can vary without any change in its weight because of difference in the density of the granulation and the pressure applied to the tablets as well as speed of tablet compression. The
thickness variation limits allowed are 5% of the size of the tablet [10, 11]. Vernier calipers were used to determine the thickness of 20 tablets. Table 2. shows mean S.D. of each brand tested.

3.3. Hardness Test. Hardness of the tablet is controlled by compression machine by applying the degree of pressure in Kg/cm² during granules compression. This parameter is very important because it effects disintegration time and dissolution profile [12]. A hardness tester (PAK/HT-0607) is used for 20 tablets randomly and mean standard deviation S. D. of each brand of cetirizine hydrochloride is calculated which shows in Table 2.

3.4. Weight variation. The weight of the tablet dosage form is measured to check the proper amount of the active ingredient in the tablet. Analytical grade weighing balance is used to measure the individual as well as average weight of the tablet and mean standard deviation S.D. of each brand of cetirizine hydrochloride is calculated which shows in Table 2.

3.5. Disintegration Time. Disintegrations are required to break up tablets into fine powder in order to increase surface area of drugs exposed to gastrointestinal fluids. Disintegration test is conducted by disintegration apparatus (IPL-27) on different brands of cetirizine 2HCl 10mg tablets and it is found that all tablets of each brand disintegrated within the range of 04-22 minutes. Figure 3 indicates that product Instazin has shown a maximum average disintegration time about 22 minutes and sample Tryzin has shown minimum disintegration time about 2 minutes which is best as compare to other products. The disintegration test do serve as a component in the overall quality control of the tablets manufacturing. The results are shown in Table 3.

3.6. Friability Test. The friability test is designed to evaluate the ability of the tablet to withstand breakage during packaging, handling, and shipping. After physical inspection and dusting, ten tablets are weighed and placed in the friability apparatus (Betatek EF-2) where they are exposed to rolling and repeated shocks as they fall 6 inches in each turn within the apparatus. After 100 revolutions, the tablets are weighed and the weight is compared with the initial weight. The loss due to abrasion is a measure of tablet friability. Maximum weight loss is not more than 1% of the initial weight of the tablet. All the results are shown in Table 2.

3.7. Thin Layer Chromatography (TLC). Thin layer chromatography is used to separate, identify and purify of the compound or drug content. The $R_f$ value of a particular pure analyte in a particular solvent (or mixture) is constant if the experimental conditions i.e. temperature, chromatography medium, solvent concentration and purity, amount of sample spotted on chromatography medium are kept unaltered. Thin layer chromatography is subject to check the specificity and purity of cetirizine hydrochloride in all brands by using solvent system ethyl acetate: methanol: formic acid (7.5:1.5:0.5 v/v/v) and compared
it by Co-TLC with the standard. It is found that the active drug in all the brands showed single spot (Rf values 0.38) after visualization on TLC plate which confirmed the purity and specificity of the drug and it was also noted that no interference of any component on TLC was observed.

3.8. Preparation of Sample and Standard Solution. An accurately weighed 10mg cetirizine 2HCl (reference standards) is transferred to 100mL volumetric flask and dissolved in 0.1N HCl and make up the volume up to the mark with the same solvent to obtain standard solution having concentration 1000ppm. Magnetic stirrer is used for better dissolution. Further 10ppm dissolution is made by taking 1mL from the above solution and make up the volume to 100mL with 0.1N HCl.

The standard solution of cetirizine 2HCl is made by dissolving 10mg of pure cetirizine 2HCl in 100mL of 0.1N HCl. Magnetic stirrer is used for better dissolution. Further 10ppm dissolution is made by taking 1mL from the above solution and make up the volume to 100mL with 0.1N HCl.

3.9. Content Assay. Every tablet contains the amount of active ingredient claimed, which can vary among the tablets within a batch. Analysis of drug potency in tablet not only indicating the presence of drug in dosage form but also shows its stability. For the evaluation of content, assay has been performed. The chemical assay results showed that all the five brands have active ingredient amount within limits i.e 90-110%.

Table 2. Physiochemical Parameter of Cetirizine 2HCl Tablets

<table>
<thead>
<tr>
<th>Brand names</th>
<th>Thickness (mm)</th>
<th>Diameter (mm)</th>
<th>Hardness test (kg/cm²)</th>
<th>Weight variation (mg)</th>
<th>Friability test (%age)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Zyrtec</td>
<td>2.71mm</td>
<td>4.10mm</td>
<td>3.08 kg/cm²</td>
<td>118 mg</td>
<td>NMT 1%</td>
</tr>
<tr>
<td>Alergo</td>
<td>3.51mm</td>
<td>5.55mm</td>
<td>3.46 kg/cm²</td>
<td>150 mg</td>
<td>NMT 1%</td>
</tr>
<tr>
<td>Cetrix</td>
<td>3.08mm</td>
<td>7.99mm</td>
<td>2.39 kg/cm²</td>
<td>160 mg</td>
<td>NMT 1%</td>
</tr>
<tr>
<td>Instazin</td>
<td>2.90mm</td>
<td>5.89mm</td>
<td>3.2 kg/cm²</td>
<td>150 mg</td>
<td>NMT 1%</td>
</tr>
<tr>
<td>Tryzin</td>
<td>3.11mm</td>
<td>4.03mm</td>
<td>3.13 kg/cm²</td>
<td>140 mg</td>
<td>NMT 1%</td>
</tr>
</tbody>
</table>

3.10. Dissolution Studies. Dissolution studies are conducted using a USP apparatus II, paddle type with 50 rpm at 371C. For standard preparation, about 10 mg of cetirizine hydrochloride is placed in a 100mL volumetric flask and dissolved with 0.1 M hydrochloric acid and then the volume was made up to
Table 3. Physiochemical Parameter of cetirizine 2HCl Tablets

<table>
<thead>
<tr>
<th>Brand names</th>
<th>Disintegration time (minutes)</th>
<th>Rf values</th>
<th>Dissolution test (%)</th>
<th>Chemical assay (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Zyrtec</td>
<td>4 minutes</td>
<td>0.38</td>
<td>89.65%</td>
<td>103%</td>
</tr>
<tr>
<td>Alergo</td>
<td>5 minutes</td>
<td>0.38</td>
<td>85.74%</td>
<td>102%</td>
</tr>
<tr>
<td>Cetrix</td>
<td>5 minutes</td>
<td>0.38</td>
<td>87.95%</td>
<td>106%</td>
</tr>
<tr>
<td>Instazin</td>
<td>22 minutes</td>
<td>0.38</td>
<td>84.25%</td>
<td>103%</td>
</tr>
<tr>
<td>Tryzin</td>
<td>2 minutes</td>
<td>0.38</td>
<td>82.70%</td>
<td>105%</td>
</tr>
</tbody>
</table>

100 mL with 0.1 M hydrochloric acid. 2 mL of this solution is transferred to another 100mL volumetric flask and diluted to 100mL with the same solvent. For the sample, about 900 mL of 0.1 mL HCl was placed in the dissolution bowl with one tablet and the apparatus was started. The sample is drawn at time intervals of 5, 10, 15, 30 and 45 minutes for each formulation. Absorbance of the sample preparation and that of standard were taken at 220 nm using a 0.1M hydrochloric acid solution as a blank. Drug concentrations are measured spectrophotometrically. Results are shown in Table 3.

4. Results and Discussion

Price fluctuation among different brands of cetirizine 2HCl is given in Figure 1. Weight variation comparison of different brands of cetirizine 2HCl is given in Figure 2. Comparison of disintegration time of all the five samples is shown in Figure 3 and comparison of % Assay of all the five samples is shown in Figure 4. It is evaluated by literatures that many drugs which are manufactured in developing countries are implicated to be substandard. In order to eliminate the health risk and provide the local community with affordable quality drugs, it is necessary to check the pharmaceutical services at regular basis. By providing this sort of information, people can make healthy choices according to their health problems. Relative to the quality of drug, the price fluctuation is also noticed in the societies where there is no regulatory control. In Figure 1 the graphical representation shows that a significant variation in the cost of different brands containing same salt i.e. cetirizine 2HCl is present, whereas there is not a significant variation present in their quality. So its use can reduce the health expenses of a patient.

A drug becomes publically famous, by its effectiveness, immediate results and lesser side effects. The effectiveness in turn depends upon the content of active ingredients, disintegration time of drug, and its dissolution. All these parameters are controlled and monitored by Pharmaceutical quality control and quality assurance department. These departments check the composition and uniformity of the drug substances used in processing and in the final product. In case of above mentioned drugs from local pharmaceuticals, the traditional tests have been used to compare them with that of a multinational company. For this
In vitro comparison of different brands

In an effort to achieve a stable and effective drug product which may be pharmaceutically equivalent to a multinational product, we conducted their physicochemical parameter’s evaluation, to get a stable and effective drug product which may be pharmaceutically equivalent to a multinational product. The safety and efficiency of these drugs was measured by uniformity content. The amount of cetirizine 2HCl in the different local brands available in Lahore, is proved to be within limits, so these can replace a multinational brand at a low cost.

The present study is based on spectrophotometric method for chemical assay, TLC technique for the purity of API’s and physiochemical parameters i.e. weight variation, hardness, friability, thickness and disintegration time as well as price fluctuation in PK rupees. Weight variation of the tablets were examined; all the tablets weight are in accordance with the required limit (NMT 5%), hardness is tested; the hardness is in good agreement with the specification (5-10 kg/cm²), thickness and the friability of the tablets are determined which complied with the specification (3.5 mm and NMT 1% ) respectively; disintegration test is performed, all the tablets were disintegrated with the prescribed time ( NMT 30 minutes); TLC technique is developed to check the quality and purity of API, all the active ingredients showed the similar $R_f$-values without any impurity (0.38) ; pharmaceutical assay was carried out, none had potency less than the required specification ( 90% to 110% ).

From the comparative study, it is observed that all the brands of cetirizine 2HCl, tablets (10mg) showed the same results as per I.P standards. Since there is a variation in the formulation of these drugs which may affect the results by performing qualitative and quantitative tests. From the test results and plots, it is observed and concluded that an economical and quality products can be prescribed for allergy patients whether they are manufactured by the local or multinational pharmaceutical companies without any health risk.

![Figure 1. Price fluctuation among different brands of cetirizine 2HCl.](image1)

![Figure 2. Weight variation comparison of different brands of cetirizine 2HCl.](image2)
Figure 3. Comparison of disintegration time of all the five samples.

Figure 4. Comparison of % Assay of all the five samples.

Competing Interests

The authors do not have any competing interests in the manuscript.

REFERENCES


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