Investigation of various practical techniques to enhance dissolution of ezetimibe from oral tablets: A comparative study

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ABSTRACT

Objective: The aim of this work was to investigate and compare various practical techniques to enhance dissolution of ezetimibe, a Class II Biopharmaceutics Classification System compound. Ezetimibe is a poorly soluble compound with variable oral bioavailability. Nanosuspensions, hydroxypropyl-beta-cyclodextrin (HPβCD) complexes as well as combination of surfactant were techniques investigated.

Materials and Methods: Nanosuspension of ezetimibe was prepared using solvent-antisolvent precipitation technique. The nanosuspensions were characterized by powder X-ray diffractometry and scanning electron microscopy. HPβCD inclusion complexes were prepared using physical mixture, co-evaporation, and kneading methods. These complexes were characterized using Fourier transform infrared spectroscopy (FTIR) and differential scanning calorimetry (DSC). Ezetimibe tablet formulations containing drug nanosuspensions, HPβCD complexes as well as sodium lauryl sulfate (SLS) enrichment were prepared and evaluated for dissolution. Results and Discussion: Nanosuspensions of ezetimibe with a mean particle size of 900 nm were successfully prepared using solvent-antisolvent precipitation. FTIR and DSC studies confirmed the formation of ezetimibe inclusion complexes. Tablets prepared using pure drug showed very poor dissolution. In contrast, tablets of nanosuspensions, HPβCD complexes and SLS mixtures showed enhanced dissolution. The tablets prepared using nanosuspensions demonstrated enhanced dissolution rate when compared to the formulations prepared using HPβCD complexes and SLS enrichment. All the techniques investigated can be used to enhance the dissolution of ezetimibe and thus can enhance the oral bioavailability and also reduce the fluctuations in the oral bioavailability. Conclusion: The techniques that were employed in this present work, nanosuspensions seem to be better when compared to cyclodextrin inclusion complexes and SLS enrichment in enhancing solubility and dissolution rate of ezetimibe.

Key words: Drug-sodium lauryl sulphate tablets, ezetimibe, hydroxypropyl-beta-cyclodextrin complexes, nanosuspension, nanosuspension tablets, sodium lauryl sulphate

INTRODUCTION

Oral drug delivery is the most common method for administration of drugs.¹ For the assessment of new drug entities, researchers and industry invariably use oral route in the exploratory stages. In the current situation, a large number of drugs reaching the hands of formulators are
poorly water soluble and thus their oral pharmacokinetics are compromised with the oral bioavailability and absorption across the gastrointestinal tract (GIT) is either low or variable. Different techniques have been successfully investigated to improve the solubility. These techniques modify the physicochemical properties of the drug or add novel functionality to the molecular structure of the drug or develop newer formulations. Such techniques include microparticles, nanoparticles, liposomes, nanosuspension, cyclodextrin inclusion complexes, surfactant-enrichment, salts, pH modification, prodrugs, nanoemulsions and re-dispersible emulsion powders. Of these many techniques, practical and clinical reality has been achieved with nanosuspensions, cyclodextrin inclusion complexes and surfactant enrichment. The enhancement in dissolution rate of a drug enhances the oral bioavailability as well as reduces the fluctuations in the bioavailability of poorly soluble compounds.

Ezetimibe is a lipid-lowering compound that selectively inhibits the absorption of cholesterol and related phytosterols from the intestine. It is a Biopharmaceutics Classification System Class II drug with low solubility and high permeability. Water solubility of ezetimibe is 8.46e-03 μg/l. It is a weakly basic compound and thus contains ionizable groups. Its log P = 4.5. Its strongest acidic pKa and strongest basic pKa are 9.48 and −3, respectively. As it is poorly soluble in water, its oral absorption is less, and it was found that it also has the problem of variable pharmacokinetics after oral administration. This could be mainly attributed to poor solubility and poor dissolution rate. Its absolute bioavailability cannot be determined as it has very low solubility that an intravenous injection cannot be prepared. Even though ezetimibe contain ionizable groups, it shows essential pH independent solubility characteristics across GIT pH range. Thus, pH-based strategies to improve solubility/dissolution characteristics (e.g. salts, addition of pH modifiers) were not the first line strategy. The other strategies can be attempted with success, but their manufacture is not practical. In this regard, nanosuspensions, cyclodextrin complexes and addition of surfactants may be the practical approaches. Thus, the objective of this study was to enhance the dissolution rate of ezetimibe so as to solve its poor and variable bioavailability problems using selected techniques, which include nanosuspensions, hydroxypropyl-beta-cyclodextrin (HPβCD) complexes and sodium lauryl sulfate (SLS) enrichment and compare each of these techniques. The techniques selected are practical and the formulations prepared using these techniques can be conveniently commercialized.

MATERIALS AND METHODS

Ezetimibe and HPβCD were obtained as a gift sample from Suven Nishtaa Ltd., Hyderabad, India. Ethanol, Tween 80 and SLS were obtained from SD fine chemicals limited. Magnesium stearate, lactose, croscarmellose sodium and povidone K 30 were obtained from Lab tech. All the other ingredients were used of analytical grade. All the equipment used in the study is routinely calibrated using procedures described by the manufacturers.

Preparation and characterization of ezetimibe nanosuspensions

Ezetimibe is a lipid lowering agent. PUMChem indicates its molecular formula as C24H21F2NO3 and molecular weight as 409.42. Ezetimibe nanosuspensions were prepared by a modification of solvent-antisolvent precipitation method as previously described for other drugs. Ethanol was used as solvent and water was used as antisolvent. Tween 80 was used as surfactant to stabilize the nanosuspension. The drug ezetimibe (100 mg) was dissolved in solvent ethanol (4 ml). A 0.2 ml of tween 80 was dissolved in distilled water (20 ml). Ethanolic solution was injected drop wise into tween 80 solution with a constant stirring on a magnetic stirrer. Ethanol was evaporated under vaccum to precipitate the nanosuspension. The product was recovered by centrifugation at 7000 rpm for 10 min using a Remi high speed centrifuge, washed twice with water and then freeze dried. The freeze dryer used was Delvac Mini Lyodel, Chennai, India. Freeze dried nanosuspension was then used for further studies. The yield of the formulation from one batch was determined using an assay method developed using ultra violet-vis. spectrophotometer. Briefly, 10 mg of ezetimibe nanosuspensions were dissolved in 10 ml of methanol, the aliquot was centrifuged and the supernatant was taken for the drug assay. The λmax used for the assay was 232 nm. The percentage yield was calculated using the formula:

\[
\% \text{ Yield} = \frac{\text{Amount of the drug in nanususpensions}}{\text{Amount of the drug taken for preparation}} \times 100
\]

The mean particle size was determined using an optical microscope. In order to examine the particle surface morphology and shape, scanning electron microscopy (SEM) was used. A concentrated aqueous suspension was spread over a slab and dried under vacuum. The sample was shadowed in a cathodic evaporator with gold layer 20 nm thick. Photographs were taken using a JSM-5200 SEM (Tokyo, Japan) operated at 20 kV. The crystallinity of the drug was evaluated using X-ray powder diffraction (XRPD). X-ray spectra were recorded with...
Preparation and characterization of hydroxypropyl-beta-cyclodextrin inclusion complexes of ezetimibe

Ezetimibe HPβCD inclusion complexes were prepared using three methods as described earlier.9,10 These methods are physical mixing, kneading method, and co-evaporation method. Preliminary phase solubility experiments were conducted to determine drug: HPβCD ratio for optimum binding. Based on the results, 1:1 weight ratio was considered optimum and we used this ratio in all the studies. In physical mixing method, ezetimibe and HPβCD were accurately weighed in equal ratios of 1:1 and were prepared by simply mixing powders with a spatula for 15 min and then sieved through 120# to obtain final desired inclusion complex. In case of kneading method, 1:1 ratio of drug and HPβCD were weighed. HPβCD and distilled water mixed together in a mortar so as to obtain a homogenous paste. Drug was added slowly; while mixing, a small quantity of methanol was added to assist the dissolution of drug. Mixtures were then blended for 2 h using an appropriate quantity of water to maintain suitable consistency. The paste was dried in oven at 45-50°C for 24 h. The dried complex was pulverized and then sieved through 120# to obtain final desired inclusion complex. In co-evaporation, drug and HPβCD were accurately weighed in the equal ratios of 1:1 and dissolved in the same quantities (5 ml) of methanol and water respectively. Both the solutions were mixed and solvents were evaporated by controlled heating at 45-50°C. The resulting solid was pulverized and then sieved through 120# to obtain final desired particle size of inclusion complex. It was made sure that the entire methanol was removed from the complex at the end of its preparation.

Ezetimibe HPβCD inclusion complexes were characterized by Fourier transform infrared spectroscopy (FTIR) and DSC. The FTIR spectra of drug, cyclodextrin, physical mixture, kneading method, and co-evaporation method were obtained. About 5 mg of sample was mixed thoroughly with 100 mg potassium bromide IR powder and compacted under vacuum a pressure of about 12 Psi for 3 min. The resulting disc was mounted in a suitable holder in Perkin Elmer IR spectrophotometer, and the IR spectrum was recorded from 4000/cm to 625/cm in a scan time of 12 min. Thermal properties of the powder samples were investigated with a differential scanning calorimetry (DSC). DSC measurements were taken by the Perkin Elmer’s PYRIS Diamond DSC. The temperature was calibrated with pure indium. All measurements were performed under a high-purity nitrogen atmosphere to minimize degradation. Approximately, 10 mg of sample was analyzed in an open aluminum pan, and heated at scanning rate of 10°C/min between 0°C and 400°C. Magnesia was used as the standard reference material.

Preparation of ezetimibe sodium lauryl sulfate powder complex

The ezetimibe SLS complex were prepared by addition of SLS in different amounts (0.1%, 0.2%, and 1%) to the drug. The powder mixture was thoroughly blended and sieved to obtain powder of desired size. This technique is called surfactant enrichment, is commonly used in industry and has been previously described.11

Preparation and evaluation of ezetimibe tablets using nanosuspensions, cyclodextrin complexes and sodium lauryl sulfate mixtures

Tablets containing pure drug, nanosuspensions, cyclodextrin inclusion complexes, and SLS enrichment were prepared using conventional wet granulation technique. Compressed tablets each containing 10 mg of ezetimibe were prepared by wet granulation method employing pure ezetimibe, nanosuspensions, cyclodextrin complexes, surfactant-enriched drug. Lactose was used as diluent to adjust the weight of the tablet to 200 mg, acacia (2%), talc (2%) and magnesium stearate (2%) were incorporated respectively as a binder and lubricants. The tablet granules were compressed into tablets on a Cadmach 16-station rotary tablet punching machine (M/s Cadmach Engineering Co. Pvt. Ltd., Mumbai, India) using 9 mm concave punches. All the tablets prepared were evaluated for content of active ingredients, hardness, friability, disintegration time and dissolution rate as per official (IP) methods. Bulk density apparatus (Cintex industrial corporation, Mumbai, India), hardness tester (Monsanto), dissolution apparatus USP (Electrolab, TDT-08 L) were used. In-vitro dissolution studies of tablet formulations were carried out using USP apparatus II paddle method. Accurately, weighed tablets were added to 900 ml of buffer media (acetate buffer pH 4.5) at 37°C ± 0.5°C and stirred at 50 rpm. An aliquot of 10 ml was withdrawn at different time intervals. An
equal volume of fresh dissolution medium was immediately replaced. The samples were assayed spectrophotometrically at 232 nm.

**Statistical analysis**

Results are expressed as mean ± standard error of the mean of six samples per treatment group. Data were analyzed using a one-way analysis of variance followed by Dunnett test. Differences were considered as significant at $P \leq 0.05$.

**RESULTS AND DISCUSSION**

In this study, various practical techniques of enhancement of solubility of a poorly soluble compound ezetimibe were investigated and compared. The novel technique that was investigated to enhance the dissolution rate of a poorly soluble compound is nanosuspensions. Poor solubility, incomplete dissolution, and insufficient efficacy are the major problems of oral drug administration. Due to smaller particle size and much larger surface to volume ratio, oral nanosuspensions or nanocrystals are specially used to solve these problems. In case of azithromycin nanosuspensions, more than 65% drug was found to be dissolved in 5 h as compared with 20% of micronized drugs. By using the standard manufacturing techniques, drug nanosuspensions can be simply incorporated into various dosage forms. The other techniques that were investigated to enhance the solubility and dissolution rate of ezetimibe are use of cyclodextrins and surfactant enrichment, which are popular in pharmaceutical formulation development.

Nanosuspension of ezetimibe was successfully prepared using the technique employed in this investigation. Particles were spherical and had an average size of 0.9 μm with a yield of 80% (Figure 1). XRPD was used to investigate the physical nature of the drug in the nanosuspensions. From the XRPD graphs, it was observed that the crystallinity of the drug was changed in the nanosuspensions (Figure 2). The peaks obtained for pure drug was very clear and sharp and the intensity of the peaks was very high when compared with peaks of ezetimibe nanosuspensions. Reduction in the peak intensity indicates the change in crystal structure. From this, we can conclude that there was reduction in the crystallinity and change into amorphous structures upon fabricating into ezetimibe nanosuspensions. Nanosuspensions of ezetimibe were not previously investigated.

Inclusion complexes of ezetimibe with HPβCD were prepared successfully by physical mixture, co-evaporation and kneading method in a weight ratio of 1:1. This was confirmed by FTIR and DSC studies. The chemical interaction between the drug and the carrier can be observed by changes in the infrared (IR) profile of complexes and can be interpreted by careful study of the spectrum. Representative FTIR spectra are shown in Figure 3. The FTIR spectra’s of the complexes were compared with spectrum of pure HPβCD and pure drug. When compared to pure drug, it has been observed that there is reduction in peak intensities of several functional groups in the complexes. Change in peak intensity indicates the complex formation. Therefore, it has been confirmed that inclusion complex of ezetimibe is formed by complexation with HPβCD. This was further corroborated by DSC studies. DSC is used to detect all processes in which energy is required or produced. The ezetimibe showed a melting peak at 184°C (Figure 4). Peak of ezetimibe at 184°C was present at the same position in all the samples with a reduction of intensity. Reduction in intensity is interpreted as formation of the complex. FTIR and DSC results suggest the trapping of drug in HPβCD.
in all the techniques. Such an interpretation was previously made in the literature. The enhancement of dissolution of ezetimibe with HPβCD was previously investigated by Taupitz et al. However, in their study, the aim was to improve the solubility of ezetimibe along with simvastatin in fixed dose combination formulations. HPβCD enhanced the dissolution rate of ezetimibe. We proved the results for ezetimibe tablets and not in combination with other drugs.

Ezetimibe tablets were prepared with the nanosuspensions, cyclodextrin inclusion complexes and surfactants using wet granulation technique. The prepared tablets were evaluated for the following parameters: weight variation, friability, hardness, disintegration time, assay, content uniformity, dissolution studies, and the results were compared. Tablet parameters indicated that good quality of tablets was prepared (Table 1). The results of dissolution studies demonstrated interesting results. The onset of dissolution of tablets prepared using unprocessed pure ezetimibe was very low (20.56% in 1 h) when compared with the tablets prepared using nanosuspensions (Figure 5). It was observed that there was an increase in % drug release (82.4% in 1 h) with tablets prepared using ezetimibe nanosuspensions.

Table 1: Evaluation of ezetimibe tablets prepared using various approaches

<table>
<thead>
<tr>
<th>Evaluation parameters</th>
<th>Nanosuspension tablets</th>
<th>Cyclodextrin complex tablets</th>
<th>Tablets containing surfactant</th>
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</thead>
<tbody>
<tr>
<td>Weight variation (%)</td>
<td>100±2</td>
<td>100±2</td>
<td>100±2</td>
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<tr>
<td>Hardness (kg/cm²)</td>
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<td>2.7</td>
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<tr>
<td>Friability (%)</td>
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<tr>
<td>Disintegration (s)</td>
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<td>15</td>
<td>15</td>
</tr>
<tr>
<td>Content uniformity (%)</td>
<td>100±0.5</td>
<td>100±0.5</td>
<td>100±0.5</td>
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Figure 3: Fourier transform infrared spectroscopy graphs of cyclodextrin complexes. (a) Pure drug; (b) cyclodextrin; (c) cyclodextrin complexes prepared using co-evaporation

Figure 4: Cumulative differential scanning calorimetry graph of drug-cyclodextrin complexes

Figure 5: Cumulative % drug release from ezetimibe tablets prepared using pure drug and nanosuspensions
It clearly indicates that nanosuspension formulation has been a successful technique to improve the dissolution rate thereby could enhance oral bioavailability and reduce fluctuations in oral bioavailability. Cyclodextrin complexes also resulted in enhancement of dissolution rate (Figure 6). Complexes prepared using co-evaporation and physical mixing significantly enhanced dissolution rates within 1 h compared with pure drug and complexes prepared using kneading technique. Inclusion complexes prepared using co-evaporation and physical mixing significantly enhanced dissolution rate of ezetimibe (59.2%, 32.75% within 1 h). Even dissolution rate of complexes prepared using kneading method also increased (28.71% within 1 h). The use of surfactants to improve the dissolution performance of poorly soluble drug products has also been successfully employed. There was an increase in dissolution as the concentration of surfactant increased (Figure 7). A 53.21% of the drug was released in 1 h in the dissolution studies with the addition of 1% SLS. All the results demonstrated statistical significance.

Although oral bioavailability of ezetimibe was not measurable because of intravenous formulation developmental problems, the studies from administering oral formulations in humans indicated that the drug exhibited moderate intersubject variability with coefficient of variation ranging from 34-43% to 32-37% for Cmax and AUC, respectively. Several formulation approaches have been addressed to reduce this variability. Such successful formulations include nanoemulsions, nanocrystals, self-nanoemulsifying granules, solid dispersions, etc. If these studies are carefully reviewed, it could indicate that nanotechnology can solve this problem of variable bioavailability of ezetimibe in a better way. Cyclodextrin inclusion complexes were also investigated to enhance the solubility and dissolution of ezetimibe and probably may reduce the fluctuations in its gastro-intestinal tract absorption. Since earlier studies have given us leads that nanotechnology can solve the problem to a better extent and since nanosuspensions of ezetimibe were not investigated, we set to investigate this approach in our study. From dissolution profiles from the tablets containing ezetimibe prepared using various techniques, nanosuspensions are found to enhance the dissolution rate to a greater extent compared with the other techniques. In this study, three different practical approaches for reduction in the variability of gastro-intestinal tract absorption of ezetimibe have been investigated, which include nanosuspensions, cyclodextrin inclusion complexes and sodium lauryl sulfate enriched formulations. All the techniques demonstrated an increase in dissolution of the drug and could therefore reduce the fluctuations in the oral bioavailability of ezetimibe. The technology can be commercialized to obtain better ezetimibe formulations.

CONCLUSIONS

Taken together, results of this work indicate that all the practical techniques employed here led to an improvement in dissolution of ezetimibe and results are comparable to similar studies conducted earlier with other techniques. Within the techniques that were employed in this research work, nanosuspensions seem to be better when compared to cyclodextrin inclusion complexes and SLS enrichment in enhancing solubility and dissolution rate of ezetimibe.

AUTHOR CONTRIBUTION

This work is a part of M. Pharm project of Prema Kumari Nannam and a part of Ph.D. project of Kiran Thadkala. Prema Kumari and Kiran have conducted the experiments.
The experiments were designed by both Dr. Chinta Sailu and Dr. Jithan Aukunuru. The manuscript was written by Dr. Jithan Aukunuru and for this editorial assistance was provided by Prema Kumari and Kiran.

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