Preparation and Characterization of Nitrendipine-loaded Eudragit RL 100 Microspheres Prepared by an Emulsion-Solvent Evaporation Method

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Abstract

Purpose: The aim of the work was to prepare nitrendipine-loaded Eudragit RL 100 microspheres to achieve sustained release nitrendipine.
Method: Nitrendipine-loaded Eudragit RL 100 microspheres were prepared by an emulsion-solvent evaporation method using ethanol/liquid paraffin system. The resultant microspheres were evaluated for average particle size, drug loading, in vitro drug release and release kinetics. FTIR spectrometry, scanning electron microscopy, differential scanning calorimetry and x-ray powder diffractometry were used to investigate the physical state of the drug in the microspheres.
Result: The mean particle size of the microspheres was influenced by varying drug:polymer ratio and emulsifier concentration while drug loading was dependent on drug:polymer ratio. The results of FTIR spectrometry, differential scanning calorimetry and x-ray diffractometry indicated the stable character of nitrendipine in drug-loaded microspheres and also revealed absence of drug-polymer interaction. The drug release profiles of the microspheres at pH 1.2 showed poor drug release characteristics while at pH 6.8, drug release was extended over a period of 8 h; release was influenced by polymer concentration and particle size. Drug release followed the Higuchi model.
Conclusion: The nitrendipine-loaded Eudragit RL 100 microspheres prepared under optimized conditions showed a good sustained release characteristics and were stable under the conditions studied.

Key words: Nitrendipine, Eudragit RL 100, Microspheres, Physicochemical characterization, Sustained release.

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INTRODUCTION
Microspheres are one of the particulate delivery systems used to achieve sustained or controlled drug delivery, improve bioavailability and stability and target drug to specific sites. Microspheres also offer advantages such as limiting fluctuation within a therapeutic range, reduction in side effects, decreased dose frequency and hence improved patient compliance. The popular method for the encapsulation of drugs within water-insoluble polymers is the emulsion solvent evaporation method. This technique offers several advantages and is preferable to other preparation methods such as spray drying, sonication and homogenization because it requires only mild conditions such as ambient temperature and constant stirring. Thus, a stable emulsion can be formed without compromising the activity of the drugs. The Eudragits are biocompatible copolymers synthesized from acrylic and methacrylic acid esters. These polymers are well tolerated by the skin and have been used in the formulation of dosage forms especially matrix tablets for oral sustained release and in tablet coating. Eudragit RL is insoluble in, but permeable to water and digestive juices, releasing drug by diffusion. Nitrendipine is an orally absorbed calcium antagonist with a short half-life of less than 3 hour; the usual oral dosage regimen is 20 mg to be taken 2 to 3 times a day. Thus nitrendipine is a suitable candidate for oral sustained drug delivery. Therefore, the present study was undertaken to prepare sustained release microspheres of nitrendipine using Eudragit RL 100 by the emulsion solvent evaporation method. The factors affecting particle size, drug loading and drug release behavior of the microspheres were investigated. The physical state of the nitrendipine was also studied using FTIR, DSC, and X-ray diffractrometry.

EXPERIMENTAL
Materials
The following materials were obtained from the indicated suppliers and used as received:

Eudragit RL 100 (Röhm GmbH & Co. KG, Darmstadt, Germany); ethanol (99%) and Span 80 (SD-Fine Chemical Works, Mumbai, India); liquid paraffin, di-sodium hydrogen phosphate anhydrous, potassium di-hydrogen phosphate, petroleum ether (40-60°C), hydrochloric acid and acetic acid glacial (Merck, Mumbai, India). nitrendipine B.P. (Concept Pharmaceuticals, Aurangabad, India)

METHODS
Preparation of nitrendipine microspheres
The nitrendipine loaded Eudragit RL100 (EURL 100) microspheres were prepared by the emulsion solvent evaporation method. In this procedure, required amount of nitrendipine and EURL 100 dissolved in ethanol, was emulsified using light liquid paraffin (80 mL) containing the emulsifier, Span 80 (2% v/v). The system was stirred continuously using a propeller stirrer at 2000 rpm and 38 ± 0.5 °C for 5 h to allow the complete evaporation of the solvent. Petroleum ether (40-60°C), 100 ml, was then added dropwise to the liquid paraffin to harden the microspheres. The paraffin was decanted off, the microspheres were washed repeatedly 4 times with petroleum ether (10 mL), collected by filtration and finally dried in a hot air oven at 40 °C for 1 h.

The nitrendipine-loaded EURL microspheres were prepared using varying polymer concentrations (1, 2 and 3% w/v) in the dispersed medium (ethanol, 20 ml) with varying drug to polymer ratios (i.e., 1:1, 1:2, 1:3, and 1:4,) while keeping the other parameter constant. The effects of process variables such as polymer concentration, drug to polymer ratio, and emulsifier concentration on the particle size of the microspheres were investigated. The optimum concentration of emulsifier was found to be 2% (v/v) and the same concentration was utilized for all the microsphere formulations.
Particle size analysis and morphological studies

The mean particle size of the nitrendipine microspheres were determined by optical microscopy. At least 200 microspheres were analyzed for each preparation and the mean diameter was calculated. The surface morphology and appearance of the microspheres were examined by a scanning electron microscopy (JEOL JSM – 5200, Japan) operating between 5-24 kV. The specimens were mounted on a metal stub (with double-sided adhesive tape) and coated under vacuum with gold in nitrogen atmosphere prior to observation. The micrographs are shown in Figure 1.

Determination of drug loading (%)

10 mg of the microspheres was dissolved in a standard flask containing absolute ethanol (10 ml) and kept for 12 hours for complete digestion at room temperature. The solution was then filtered through a filter disc (particle
retention: 11µm) and the filtrate was assayed spectrophotometrically for drug content at 356 nm to compute the drug concentration from the standard curve. The drug loading (%) of the microspheres was calculated using the equations given below.

\[ \text{Drug loading} (\%) = \frac{M_{\text{actual}}}{\text{weight of powdered microspheres}} \times 100 \]

where \( M_{\text{actual}} \) is the actual drug content in sample of microspheres.

**In vitro drug release studies**

Drug-release tests on the microspheres were carried out using a USP dissolution rate test apparatus Type II (Electro Lab model TDT-08L) for 8 h at a stirring speed of 100 rpm and temperature of 37 ± 0.5 °C. An amount of the microspheres equivalent to 20 mg of nitrendipine filled in a hard gelatin capsule (Size no.2) were placed in the dissolution medium containing 600 ml of phosphate buffer (pH 6.8) B.P. containing 0.3 % (w/v) of sodium dodecyl sulphate (SDS) to maintain sink condition for the drug. A quantity (5ml) of the dissolution medium was sampled at predetermined time intervals, and fresh dissolution medium was simultaneously used replenish the dissolution medium on each occasion to keep the volume constant. The sample was filtered through filter disc (particle retention: 11µm), the filtrate diluted to 10 ml with fresh dissolution medium and assayed spectrophotometrically (160 – UV - visible Shimadzu spectrophotometer) at 356 nm to determine drug concentration. The same procedure was used to assess drug release pattern in hydrochloric acid (0.1M) buffer solution (pH 1.2). The release kinetics of nitrendipine from the microspheres was determined using different models, viz, zero order, First order, Higuchi model, Peppas-Korsmeyer and Hixon-Crowell.

**FTIR spectroscopic studies**

The IR spectra of the samples were recorded on an FTIR spectrophotometer (Perkin Elmer 1600 series) using KBr pellet (12 mm disc), compressed in a hydraulic press at 10 tons for 30 seconds.

**Differential scanning calorimetric analysis**

Differential scanning calorimetry (DSC) analysis was undertaken to characterize the changes, if any, during thermal exposure of samples. The test was carried out using a thermal analysis system (Mettler TA 4000 model). Calibration with the standard (indium) was undertaken prior to subjecting the samples for study (between 30-400 °C), which were heated at 10° C/min in an aluminum pan under a nitrogen atmosphere while using an empty pan as the reference in this instrument. The instrument automatically calculated onsets of melting point and enthalpy of fusion. The results are shown in Figure 4.

**X-ray powder diffraction analysis**

The x-ray diffraction pattern of the samples were obtained using an x-ray diffractometer (Rich Seifert, model 3000 P) at 30 kV, 15 mA over a range of 10-100 2θ, using Cu Kα radiation wavelength of 1.5405 Å. In this technique, the cavity of the metal sample holder of the x-ray diffractometer was filled with ground sample powder and then smoothed out with a spatula. The results are shown in Figure 5.

**RESULTS**

**Particle size and morphological characteristics**

Figure 1 shows the scanning electron photomicrograph of the surface of nitrendipine-loaded EURL microspheres having mean particle size 42.15 ± 12.34 µm. The particle size of the microspheres increased from 14.25 ± 6.85 to 42.15 ± 12.34 µm with increasing polymer concentration (Table 1). Few drug crystals appeared on the surface of the microspheres. It was also observed that increasing the emulsifier concentration decreased the microsphere size.

**Drug loading of the microspheres**

Increase in the concentration of drug in the organic solvent resulted in an increase in the drug loading of the microspheres. The results indicate that the highest nitrendipine loading
of 38.40 ± 0.44 % was observed when the polymer concentration was 3% at a drug: polymer ratio of 1:1 and a stirring rate of 2000 rpm.

**In vitro** drug release and release kinetic model studies

Figure 2 indicate that maximum drug release from nitrendipine-loaded microspheres at pH 1.2 was about 40% over a period of 4 h.
Furthermore, there was no significant difference in drug release characteristics of the EURL microspheres irrespective of the polymer concentration and drug: polymer ratio.

Figure 3 illustrates the cumulative percent release of nitrendipine-loaded microspheres at pH 6.8 over a period of 8 h. The data clearly show that drug release (for microspheres prepared at a drug polymer ratio of 1:1) decreased with increase in the polymer concentration as follows: 1, 2, and 3 % polymer concentrations showed 93.69 ± 0.95%, 89.37 ± 1.01%, 82.65 ± 0.45% drug release, respectively. At other drug : polymer ratios (i.e., 1:2, 1:3, and 1:4) there was only a small retardation of drug release from the microspheres ranging from 86.57 ± 0.84% to 73.40 ± 0.79% (see Figure 3). The EURL microspheres prepared at 3% polymer concentration showed a certain level of sustained release characteristics, especially microspheres prepared with 1:3 drug to polymer ratio (E3d) which released 73.40 ± 0.79% of drug at pH 6.8 over 8 h. An initial 20-33 % drug release was observed in all the microsphere formulations within the first hour at pH 6.8. Based on correlation coefficient ($r^2$) data, the release pattern of nitrendipine from Eudragit RL microspheres fitted best to the Higuchi model.

**FTIR- spectroscopic studies**

FTIR of nitrendipine shows the principal peaks at the wave numbers of 1020.66 – 1349.17 cm$^{-1}$, indicating the presence of carboxyl, carboxylate groups, and carbonyl ester vibrations at 1701.64 cm$^{-1}$, while C-H stretching appeared at 2991.62 – 2902.96 cm$^{-1}$ and NO$_2$ group appeared between 1349.17 and 1531.18 cm$^{-1}$. In the IR spectra of the physical mixture of the formulation ingredients as well as those of nitrendipine-loaded EURL 100 microspheres (E3d), the principal peaks for nitrendipine in the formulation, (E3d), appeared between 1020.66 and 1349.56 cm$^{-1}$ and indicates the presence of carboxyl, carboxylate groups and carbonyl ester vibration at 1701.74 cm$^{-1}$, as well as NO$_2$ group between 1349.56 and 1532.34 cm$^{-1}$. Therefore, the possibility of any drug polymer interaction is ruled out.

**Differential scanning calorimetric studies**

Figure 4 illustrates the DSC thermogram of nitrendipine, physical mixture of nitrendipine with EURL, and nitrendipine-loaded EURL 100 microspheres (E3d). The DSC thermograms of the physical mixture of nitrendipine with Eudragit RL, on the one hand, and the microsphere formulation (E3d) on the other, produced almost similar melting endotherms of pure drug at 158.6 °C and 156.0 3 °C, respectively. However, the intensity of the drug fusion peak for the microsphere formulation was lower than that of the pure drug and physical mixture.

**X-ray diffraction studies**

Figure 5 illustrates the comparative X-ray powder diffraction pattern of nitrendipine alone, physical mixture of nitrendipine with EURL, and nitrendipine-loaded EURL microspheres (E3d). The x-ray diffraction profile of Eudragit RL polymer indicated the presence of a completely amorphous material; pure nitrendipine showed the classical diffractogram of the crystalline product. No major difference in the XRD patterns of the physical mixture and the drug loaded microspheres was noticed. However, decreases in the peak intensity and the baseline shift of the diffractogram were observed in the case of the nitrendipine-loaded microspheres (E3d) when compared to that of the physical mixture. This is possibly due to the decrease in the degree of crystalinity of the drug following dispersal in the polymer matrix.

**DISCUSSION**

Nitrendipine-loaded Eudragit RL microspheres can be easily prepared by solvent evaporation method; the microspheres were satisfactory considering their size and shape. The microsphere size increased with increasing polymer concentration; this may be due to increasing polymer concentration that produced a significant increase in the viscosity, thus leading to an increase of the emulsion droplet size and finally a larger
microspheres size. On the other hand, microsphere size decreased with increasing emulsifier concentration. The effect of emulsifier concentration on the size of the microspheres can be explained by the higher stability of the dispersed droplets and less tendency of the droplets to coalesce upon collision at higher concentrations of the surfactant. The appearance of drug crystals on the surface of the microspheres may be attributed to high drug concentrations and

Figure 4: DSC thermograms of Eudragit RL100 (a); Eudragit RL100 microspheres without drug (b); nitrendipine (c); physical mixture of nitrendipine and Eudragit RL100 (d); and nitrendipine-loaded Eudragit RL100 microspheres (e).

Figure 5: X-ray powder diffraction patterns of Eudragit RL100 (a); Eudragit RL100 microspheres without drug (b); nitrendipine (c); physical mixture of nitrendipine and Eudragit RL100 (d); and nitrendipine-loaded Eudragit RL100 microspheres (e).
slow solvent removal as the drug formed a particulate (crystal) dispersion resulting. Drug loading increased with increase in the drug proportion of the preparation. The results of in vitro drug release studies showed that the various drug: polymer ratios and different polymer concentrations had a considerable effect on drug release pattern. The poor drug release and insignificant variation in the drug release pattern of nitrendipine-loaded microspheres at pH 1.2 may be attributed to the effect of the dissolution medium on drug dissolution from the microspheres. The microspheres prepared at a polymer concentration 3% showed considerable sustained release characteristics, especially microspheres prepared at a 1:3 drug: polymer ratio (E3d) which released 73.40 ± 0.79% of drug at pH 6.8 over a period of 8 h and showed better sustained release characteristics when compared with other microsphere types (Figure 3). This may be attributed to the higher polymer content which resulted in a larger particle size and a tightened polymer network and thus retarding drug release. The initial 20-33% of drug release from all the microspheres may be attributed to the presence of a small proportion of the drug on the microsphere surface which could have arisen from the diffusion of the drug during preparation and drying. When such microspheres are immersed in dissolution medium, the surface drug is immediately released. The assessment of the release kinetics revealed that drug release from the microspheres at pH 6.8 phosphate buffer followed the Higuchi model. The DSC thermographs of the microspheres showed that the drug fusion peak intensity was less than those for the pure drug and the physical mixture. This may be due to the reduced drug content, (arising from homogeneous dissolution of the drug in the polymer. Other tests indicate that nitrendipine remained in a crystalline state with in the polymer network of the microsphere. It is clearly evident from the XRD pattern, however, that lower peak intensity and a baseline shift were observed for the microsphere when compared to that of the physical mixture. This may be due to a decrease in the degree of crystallinity of the drug.

CONCLUSION

From the foregoing investigations it was concluded that the nitrendipine loaded Eudragit RL microspheres under optimized conditions showed some degree sustained release and were stable under the conditions studied. The release kinetics followed the Higuchi model.

REFERENCES