Design and *in vitro* evaluation of Eudragit-based extended release diltiazem microspheres for once and twice daily administrations: the effect of coating on drug release behavior

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**Short title:** Extended release diltiazem microspheres
ABSTRACT

Objectives: The aim of this investigation was to develop an extended release formulation of diltiazem hydrochloride (DL) for once and twice-daily administration, based on Eudragit (Eud) RL and RS microspheres using emulsion solvent evaporation method.

Methods: Formulations with different drug-polymer concentrations were produced and characterized in terms of yield, encapsulation efficiency (EE), particle size and surface morphology. The drug release and thermal behavior of microspheres were also investigated. Selected microspheres were then coated with Eud RS by continuous solvent evaporation method, in order to modify the microspheres properties and burst release.

Results: According to the results, the EE was in the range of 56-93 % for uncoated microspheres. The mean particle size of microspheres was different from 470 to above 1000 µm, based on various formulation variables. No difference was observed between the mean size of particles prepared with Eud RL and Eud RS. Microspheres showed sustained release behavior which was affected by drug: polymer ratio as well as particle size. Coating the microspheres not only improved the EE values (82-92 %), but also reduced the mean dissolution rate (MDR) and also the burst release.

Conclusion: Microspheres prepared with DL: Eud RL ratios of 1:3 and 1:4, showed the release profiles in accordance with the USP criteria for DL extended release product, for dosing every 12 and 24 h, respectively.

Key words: coating, diltiazem hydrochloride, eudragit RL and RS, extended release, microspheres.
INTRODUCTION

Diltiazem hydrochloride (DL) is a highly soluble calcium channel blocker drug which is used in the treatment of high blood pressure and angina pectoris.\textsuperscript{1} Due to its short elimination half-life of 2-5 hours, the conventional oral dosage forms are administered 3-4 times a day, to maintain an effective plasma concentration, which resulted in low and variable bioavailability.\textsuperscript{2} Using sustained release form of this medication is vital for its efficacy by achieving relative constant blood concentrations, improving the clinical efficacy of drug as well as patient compliance.

However, along with the benefits of using extended release single-unit tablets, there are some limitations for these systems, such as dose adjustment problem and the effect of food on drug release. Besides, breaking those tablets before taking the medicine could cause a different release behavior and serious side effects.\textsuperscript{3} The above-mentioned problems could be overcome using microspheres as multiple-unit dosage forms. Microspheres uniformly distribute in the gastrointestinal tract and resulted in more uniform drug absorption, limited fluctuation within a therapeutic range, decreased dose frequency and reduced patient-to-patient variability.\textsuperscript{4}

The physical properties and release behavior of microspheres are dependent on different factors such as drug and polymer nature as well as method of manufacturing. According to the literatures, the existence of drug particles on the surface, particles embedded in the surface layers as well as high porosity of microspheres are considered as main reasons for initial burst release. Coating of microspheres is one of the approaches to reduce the burst release and modify the drug release behavior.\textsuperscript{5} Preparation and separation of initial microspheres and then using them in a separate coating process would be time and cost consuming. Therefore application of a continuous preparation and coating process of microspheres seems to be more preferable.

To date, modified release microspheres of DL using different polymers and methods was developed in order to extend its clinical effects.\textsuperscript{2, 6-11} Only a few researches were performed on the preparation of Eud RS-based DL microspheres by solvent evaporation method.\textsuperscript{12-14}
The objective of the present research was to design and evaluate DL-loaded Eud RL and RS matrix-type microspheres as extended release systems for both once and twice-daily administrations, in order to reduce its dosing frequency. With the aim of achieving both systems, different drug-polymer concentrations were applied and examined. In addition, the effect of coating of microspheres by Eud RS on drug release behavior and the burst effect was also evaluated. Emulsion solvent evaporation method was used for microspheres preparation as well as continuous coating process. This is simple method that has been used to prepare microspheres of different soluble and insoluble compounds.  

MATERIALS AND METHODS

Materials

Diltiazem Hydrochloride (DL) powder (Zambon Group SPA, Italy), Eudragit (Eud) RL and RS 100 (Röhm Pharma GmbH, Germany), span 60 (Sigma-Aldrich, St. Louis, MO, USA), n-hexane (Carlo Erba, France) and liquid paraffin (Merck, Germany) were used in this study. Materials and excipients used in preparing microspheres were of pharmacopoeial grades.

Microspheres preparation

DL-loaded Eud RL and RS microspheres were produced by emulsion solvent evaporation method. Different amount of drug and polymer were dissolved in 3 ml ethanol (dispersed phase) which was then slowly (with the rate of 1 ml/min) added to a beaker containing a mixture of 50 ml liquid paraffin and 0.1 % w/v span 60 (continuous phase) while stirring at the speed of 500 rpm under a mechanical stirrer (IKA, Germany). The mixture was stirred until the organic solvent evaporated completely. The prepared microspheres were collected by filtration and washed three times with n-hexane until the whole paraffin was removed. Finally, the microspheres were dried at room temperature for 24 h and kept in air tight containers for further studies.

Coating of microspheres

A one-step continuous solvent evaporation technique was used coating process. Primary microspheres were prepared by the above-mentioned method, but before
completing the process and collecting the microspheres, a 3.3 % w/v Eud RS ethanolic solution was added drop wise to the continuous phase and stirred until complete solvent evaporation. The other steps were similar to the previous method.

Characterization of microspheres

The prepared microspheres were characterized in terms of yield value, encapsulation efficiency (EE), morphology, drug release, particle size and thermal analysis. The yield value of each formulation was calculated by the following equation: yield value (%) = (weight of dried microspheres / total solid material amount in the dispersed phase) ×100.

Drug content

Ten mg of dried microspheres was accurately weighted and transferred to a beaker containing 10 ml methanol and stirred for 15 min to dissolve microspheres completely. The solution was analyzed for DL content by a UV spectrophotometer (Shimadzu UV1201, Japan) at 240 nm after dilution. The drug loading and encapsulation efficiency (EE) were calculated using the following equations:

Drug loading (%) = (weight of drug in microspheres/weight of microspheres) ×100
Drug EE (%) = (actual drug loading/theoretical drug loading) ×100.

In vitro drug release

Drug release of all microspheres was carried out using USP type II dissolution test apparatus (Erweka DT6R, Germany) in 900 ml phosphate buffer solution (pH=7.2) at 37 ± 0.5°C at a rotation speed of 50 rpm (in accordance with the USP test number 5 for DL extended release form dosing every 12 h). Three ml of the medium was withdrawn at predetermined time intervals and replaced with the same amount of fresh dissolution medium after each sampling. The sample solutions were analyzed for drug content at 240 nm by a UV spectrophotometer.

The dissolution test was also performed on selected microspheres in compliance with the USP test number 2 for DL extended release form dosing every 24 h, using an apparatus II at 100 rpm and 900 ml of dissolution medium (distilled water) for 15 h. All experiments were performed in triplicate for each formulation.
All formulations were compared using different dissolution parameters. Mean dissolution time (MDT) which was applied to analyse dissolution profiles was calculated arithmetically by the following equation:

$$\text{MDT} = \frac{\sum_{i=1}^{n} t_i \Delta M_i}{\sum_{i=1}^{n} \Delta M_i}$$

where $\Delta M_i$ is the fraction of drug released in time $t_i$ (calculated by $(t_i + t_{i-1})/2$) and $i$ is the sample number.

In addition, the area under the dissolution curve (dissolution efficiency; DE) was calculated by the below formula:

$$\text{DE} = \frac{\int_{t_0}^{t_{100}} y \, dt}{y_{100} \times 100}$$

Where $y$ is the percentage of drug dissolved at time $t$. Mean dissolution rate (MDR) was also calculated based on the following equation:

$$\text{MDR} = \frac{\sum_{i=1}^{n} \Delta M_i/\Delta t}{n}$$

where $\Delta t$ is the time at the midpoint between $t$ and $t-1$ and $n$ is the number of dissolution sample times.

**Particle size**

The mean particle size of DL microspheres was determined by optical microscopy. At least 200 microspheres were analyzed for each preparation and the mean diameter was calculated.

**Surface morphology**

The appearance and surface morphology of microspheres were evaluated by scanning electron microscopy (SEM, Philips XL30, The Netherlands). Microspheres were attached to a specimen holder with double-sided adhesive tape and coated under vacuum by gold sputter coater (Bal-Tec SCD 005, Switzerland) prior to observation.

**Differential Scanning Calorimetry (DSC)**

DSC analysis of drug, polymer, selected DL-loaded microspheres and related physical mixture was done. After calibrating the apparatus (Shimadzu DSC-60, Japan) by indium standard, accurately weighed samples (5 mg) were placed in sealed aluminum pans. The containers were placed in DSC apparatus and heated at a constant rate of 10/min.
over a temperature range of 25 to 300 °C. An empty standard aluminum pan was used as reference.

**Statistical analysis**

Statistical analysis of different variables was carried out using analysis of variance (ANOVA) followed by Tukey’s post test. Significance was tested at the 0.05 level of probability.

**RESULTS AND DISCUSSION**

DL microspheres were successfully prepared by emulsion solvent evaporation method using ethanol as drug-polymer solvent (dispersed phase) and liquid paraffin-span 60 mixture as continuous phase. The yield value was in the range of 62.8-92.4 % for initial microspheres and 81.3-97.6 % for Eud RS-coated microparticles.

**Characterization of microspheres**

*Encapsulation efficiency and particle size*

Table 1 shows the composition and properties of the Eud RL and RS-based microspheres prepared with different drug-polymer ratios. Increasing the amount of Eud RL from 300-800 mg, lead to a 25 % enhancement in the EE values. In fact, the size of emulsion droplets was increased due to the higher viscosity of the polymeric solution, which in turn decreased the surface area and also drug molecule transport from dispersed to continuous phases. Particle size of those microspheres was also increased significantly (P< 0.001) which was expected. The similar results were obtained for the microspheres prepared with higher DL concentrations. Based on the results (Table 1), there is a significant difference between the EE values of M8L and M4L (P< 0.05). In addition, using higher drug: polymer ratios resulted in significantly (P< 0.001) increased particle size. Although the effect of polymer concentration on the particle size seemed to be more than drug, but the results revealed that in certain drug concentrations, its effect on particle size cannot be neglected.

Application of various drug-polymer concentrations with the same ratio (M1L, M2L and M7L) resulted in microspheres with different EEs and mean particle sizes. An increase of 20 % was found for the EE value of M7L (higher DL- Eud RL concentration)
compared to M1L and M2L. In other words, an appropriate increase in drug and polymer concentrations simultaneously, led to more drug entrapment in microspheres. The same trend was also observed for microparticles size which could be attributed to the higher viscosity and emulsion droplet size of this formulation. Although the difference observed between M2L and M1L was far less.

By changing the polymer type from Eud RL to Eud RS (Table 1), no difference was noticed in EE % for lower drug: polymer concentrations (M1L and M1S). However, this was opposite for the formulations prepared with higher drug: polymer concentrations, in which M2S showed improved EE value compared to M2L, which is in accordance with some literatures. Eud RL is more permeable and the diffusion of drug molecules from the droplets to the surrounding medium during preparation process is more probable than Eud RS. In addition, the repulsion between quaternary ammonium groups of Eud RL and cationic drug could facilitate the DL removal to the external phase and reduced the EE.

According to the results, coating of microspheres improved the EE % significantly (P<0.001) compared to the uncoated microspheres (Table 2). It is probable that application of Eud RS coating on the surface of initial microspheres prevents the drug molecules transport to the emulsion external phase during preparation process. Meanwhile, no difference was observed in the EE values of the coated microspheres with different inner polymers. As it was expected, the mean particle size of microparticles was increased following coating process. The higher mean particle size of M2LS and M2SS compare to the M1LS and M1SS was related to the higher inner polymer concentration used to prepare the initial microspheres.

**SEM**

The SEM micrographs (Figure 1) show that microspheres prepared in the presence of lower polymer concentration (M2L) were more spherical in shape with wrinkled surfaces compared to M5L (higher polymer amount). Using higher DL concentration in the formulations, did not affect the microspheres shape, but increased their roughness mainly due to the existence of drug crystals on the surface layers of microspheres. No difference was observed between the microspheres prepared by Eud RL and RS (M1L
and M1S) in terms of shape and surface properties which was in accordance with a previous research.24 Based on the results, following coating of microspheres, they were still spherical in shape with more uniform surfaces compare to their initial uncoated microparticles. The study of surface morphology of M1LS and M1SS (Figure 1) confirmed the absence of drug crystals on the surface of microparticles and suitable coverage of the initial microspheres during the continuous coating process.

**Drug release studies**

The release profiles of DL from microspheres prepared with different formulations was presented in Figure 2. Based on the results, the drug release rate was decreased apparently with increasing the polymer concentration (Figure 2A). The DL released after 3 hours of the experiment for M2L and M5L was 73.25 and 26.09 %, respectively. This trend could be also observed in MDR and DE values (Table 1). Besides, decreasing the drug release rate led to an increase in MDT values. In fact, higher polymer concentration resulted in larger particle size with less surface area and therefore lower release rate. A burst release of about 37 % was observed for M2L during the first hour of the study which could be attributed to the lower polymer content and particle size, as well as more drug particles exist on the surface layers of microspheres.

Using higher drug concentrations with the fixed amount of the polymer enhanced the drug release apparently (Figure 2B) and about 90 % of DL was dissolved during the 5 hours from M8L (DE=80.78 ± 1.91 %). Table 1 shows that MDR was significantly increased in formulations prepared with higher drug concentrations (P< 0.0001). In addition, the burst release of these microspheres was in the range of 11.88-42.70 %. It seems that the presence of more drug particles on the surface layers of microspheres prepared with higher drug levels, enhanced release drug rate in spite of larger particle size.15 In fact, reduction of the drug diffusion pathway is possible in microspheres with higher drug loading.25 Also removal of drug particles from microspheres leads to the formation of more porous structure that plays an important role in accelerating drug release.26

Using various drug: polymer concentrations with the same ratio also led to microspheres with different release behavior. Based on Table 1, M7L prepared with higher drug-
polymer concentration, extended the drug release more than M2L and M1L (P < 0.001), mainly due to its larger particle size. Significant decrease in MDR value for M7L (P < 0.001) corresponds to an increase of more than 110 min in MDT of this formulation in comparison to M1L. All those three microspheres showed a burst release in the range of 30-50 %.

Figure 2C shows the release profiles of DL from microspheres prepared with Eud RL and RS. It is obvious that the drug release from Eud RS-based microspheres was slower than particles made with Eud RL. The difference observed between M1L and M1S was more evident. Based on Table 1, a reduction of more than 18 % in DE and about 1.5 folds in MDR was observed for M1S compared to M1L. The MDT values for M1S and M2S were also significantly more than M1L and M2L (P < 0.001). Since the mean particle size was not very much affected by the polymer type, the obtained results could be attributed to the lower permeability of Eud RS.

The effect of coating of microspheres on DL release profile was illustrated in Figures 2D and 2E. The drug release from all coated particles was decreased apparently compared to the uncoated microspheres. Based on Table 2, a significant reduction in MDR and DE values was observed for coated particles. The lowest MDR belongs to the formulation M2SS, which was about ½ times lower than that of M2S. The lowest DE % was also obtained for M2SS. Decreasing the release rate was along with increasing the MDTs. A significant difference was observed between the MDTs of M1L and M1LS and also M2L and M2LS (P < 0.01). Following coating process, MDT of the microspheres with Eud RL as inner polymer enhanced more than that of Eud RS. Furthermore, although the burst release was declined for all coated microspheres, but this was more noticeably for microspheres with Eud RL as core polymer.

The results revealed that although coating of microspheres was helpful in decreasing the drug release rate, but it was not as effective as using appropriate drug-polymer concentration in the preparation process, without any coating. Formulations M5L and M6L showed the least MDRs and the highest MDT values among all coated and uncoated microspheres. It seems that the drug particles in the mass of microspheres was much more than the particles in the surface layers and controlling their diffusion was more important in achieving the desirable extended release behavior. However,
application of higher polymer concentration in the coating process could show different effects.

**DL microspheres for once and twice-daily administrations**

Figure 2F shows the release profiles of two selected formulations (M4L and M5L) in phosphate buffer solution (pH= 7.2) (USP test number 5) and water (USP test number 2), respectively. The results indicated that microspheres prepared with DL: Eud RL ratios of 1:3 (M4L) and 1:4 (M5L) were in accordance with the USP test for DL extended release form dosing every 12 and 24 h, respectively, without any further treatment.

The release kinetics of these formulations was investigated using three different models including zero order, first order and Higuchi equation. Based on the squared correlation coefficient ($R^2$), the release profile of M4L was best fitted with zero order ($R^2=0.989$) compared with first order and Higuchi model ($R^2 = 0.973$ and 0.944, respectively). Although the $R^2$ values calculated for M5L based on first order (0.991) and Higuchi equation (0.994) were higher than that of zero order (0.954), but there is no evidence to specify the dominant kinetics model for this formulation.

**DSC**

The DSC thermograms of DL, Eud RL, selected microsphere (M4L) and related physical mixture (PM) were depicted in Figure 3. A characteristic endotherm appeared for the drug at the onset temperature of 210.08 °C which could be attributed to the melting of DL. A broad peak in the range of 50-60 °C was observed in the thermogram of Eud RL which is related to its glass transition temperature. The DSC curve obtained for the microspheres presented the same thermal profile as that of physical mixture, both containing the drug melting peak with a slight shift toward lower temperatures. These minor changes in the drug endotherm could be attributed to the presence of polymer, which lowers the drug purity. This result suggests no interaction between the drug and the polymer during the preparation process.

**CONCLUSION**

DL: Eud RL extended release microspheres for once and twice-daily administrations for the treatment of hypertension and angina pectoris were successfully produced in this study by a simple method of solvent evaporation using suitable formulation variables.
The results confirmed that a one-step continuous emulsion solvent evaporation process was a practical technique to prepare coated microspheres with improved physical properties (especially EE %) and reduced burst release. Using suitable drug: polymer ratios as well as external coating polymer concentration could modify the particle size, surface morphology, porosity and the amount of drug particles on the surface layers which are essential to obtain desirable results.

REFERENCES


Table 1: Composition and physicochemical properties of DL microspheres (mean ± SD, n=3)

<table>
<thead>
<tr>
<th>Formulation</th>
<th>Drug (mg)</th>
<th>Polymer (mg)</th>
<th>EE(^c) (%)</th>
<th>Mean particle size (µm)</th>
<th>MDT(^d) (min)</th>
<th>DE(^e) (%)</th>
<th>MDR(^f) (%min(^{-1}))</th>
</tr>
</thead>
<tbody>
<tr>
<td>M1L(^a)</td>
<td>100</td>
<td>150</td>
<td>61.43±1.33</td>
<td>452.9±7.29</td>
<td>82.45±4.17</td>
<td>83.78±0.98</td>
<td>0.331±0.002</td>
</tr>
<tr>
<td>M2L</td>
<td>200</td>
<td>300</td>
<td>62.15±1.09</td>
<td>513.8±10.09</td>
<td>94.38±4.05</td>
<td>75.29±0.58</td>
<td>0.210±0.004</td>
</tr>
<tr>
<td>M3L</td>
<td>200</td>
<td>500</td>
<td>56.62±3.75</td>
<td>620.0±5.82</td>
<td>150.70±6.77</td>
<td>74.96±0.54</td>
<td>0.174±0.007</td>
</tr>
<tr>
<td>M4L</td>
<td>200</td>
<td>600</td>
<td>81.46±2.60</td>
<td>665.7±6.71</td>
<td>197.11±5.62</td>
<td>61.43±1.11</td>
<td>0.086±0.004</td>
</tr>
<tr>
<td>M5L</td>
<td>200</td>
<td>800</td>
<td>87.70±3.57</td>
<td>720.1±12.58</td>
<td>210.27±2.34</td>
<td>54.00±1.79</td>
<td>0.068±0.006</td>
</tr>
<tr>
<td>M6L</td>
<td>300</td>
<td>600</td>
<td>82.18±0.84</td>
<td>745.9±5.58</td>
<td>235.95±7.86</td>
<td>58.29±1.09</td>
<td>0.064±0.002</td>
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<tr>
<td>M7L</td>
<td>400</td>
<td>600</td>
<td>84.97±3.53</td>
<td>813.2±2.33</td>
<td>201.50±4.26</td>
<td>66.98±0.43</td>
<td>0.147±0.005</td>
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<tr>
<td>M8L</td>
<td>500</td>
<td>600</td>
<td>92.86±3.90</td>
<td>1027.3±6.50</td>
<td>114.94±6.45</td>
<td>80.78±1.91</td>
<td>0.246±0.007</td>
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<tr>
<td>M1S(^b)</td>
<td>100</td>
<td>150</td>
<td>59.53±1.19</td>
<td>463.5±4.05</td>
<td>116.74±1.05</td>
<td>65.00±1.93</td>
<td>0.211±0.005</td>
</tr>
<tr>
<td>M2S</td>
<td>200</td>
<td>300</td>
<td>77.09±1.05</td>
<td>528.6±3.35</td>
<td>102.57±1.71</td>
<td>63.95±0.36</td>
<td>0.224±0.011</td>
</tr>
</tbody>
</table>

\(^a\) L: Eud RL, \(^b\) S: Eud RS, \(^c\) EE: Encapsulation efficiency, \(^d\) MDT: Mean dissolution time, \(^e\) DE: Dissolution efficiency, \(^f\) MDR: mean dissolution rate
**Table 2:** Composition and physicochemical properties of DL microspheres coated with Eud RS (mean ± SD, n=3)

<table>
<thead>
<tr>
<th>Formulation</th>
<th>Core drug (mg)</th>
<th>Core polymer (mg)</th>
<th>Type of core polymer</th>
<th>EE a (%)</th>
<th>Mean particle size (µm)</th>
<th>MDT b (min)</th>
<th>DE c (%)</th>
<th>MDR d (%/min)</th>
</tr>
</thead>
<tbody>
<tr>
<td>M1LS</td>
<td>100</td>
<td>150</td>
<td>Eud RL</td>
<td>91.99 ± 0.82</td>
<td>510.4 ± 5.85</td>
<td>118.04 ± 3.27</td>
<td>77.42 ± 0.59</td>
<td>0.195 ± 0.004</td>
</tr>
<tr>
<td>M1SS</td>
<td>100</td>
<td>150</td>
<td>Eud RS</td>
<td>88.32 ± 1.31</td>
<td>500.2 ± 3.05</td>
<td>131.16 ± 5.71</td>
<td>57.25 ± 1.48</td>
<td>0.184 ± 0.007</td>
</tr>
<tr>
<td>M2LS</td>
<td>200</td>
<td>300</td>
<td>Eud RL</td>
<td>82.08 ± 2.05</td>
<td>641.1 ± 7.81</td>
<td>125.18 ± 5.32</td>
<td>68.92 ± 1.53</td>
<td>0.146 ± 0.004</td>
</tr>
<tr>
<td>M2SS</td>
<td>200</td>
<td>300</td>
<td>Eud RS</td>
<td>83.84 ± 1.92</td>
<td>610.5 ± 3.52</td>
<td>138.87 ± 2.30</td>
<td>56.83 ± 0.20</td>
<td>0.106 ± 0.002</td>
</tr>
</tbody>
</table>

a EE: Encapsulation efficiency, b MDT: Mean dissolution time, c DE: Dissolution efficiency, d MDR: mean dissolution rate
Figure 1: SEM micrographs of different microspheres and their surfaces
Figure 2: Release profiles of DL from A, B and C: microspheres with different formulation variables in phosphate buffer (pH=7.2), D and E: coated versus uncoated microspheres, F: M4L in phosphate buffer (pH=7.2) and M5L in water (n=3)
Figure 3: DSC thermograms of DL, Eud RL, M4L and related physical mixture (PM)