Establishment and Escalation of Amino Acid Stacked Repressible Release Embedded System Using QbD

QbD'yi Kullanarak Amino Asit Yığılmış Bastırılabilir Salınan Gömülü Sistemin Kurulması ve Arttırılması

Abstract

Objectives: Traditional approach of developing a new delivery system is an exhaustive task and requires a number of resources like man, money, material and time. To overcome this problem Quality by Design (QbD) can be utilized to get the pharmaceutical product of desired (best) quality with minimum use of above resources as well as determination of impact of one factor over the desired associated process. The present research is focused on establishing a design for formulating optimized gelatin microspheres using QbD.

Materials and Methods: Characterization of formulated microspheres was done by infra red spectroscopy, scanning electron microscopy, percentage yield, microsphere size, drug entrapment efficiency and drug release. Impact of concentration of gelatin and ethyl cellulose was determine over dependent response like percentage yield, microsphere size and drug entrapment efficiency.

Results: Response surface curve was obtained by using $3^2$ central composite design and optimized batch was obtained with percentage yield, microsphere size and drug entrapment efficiency as 89.98, 333.32 mm and 82.61% respectively. Validation of optimized batch was done formulating four different batches with optimized values of independent response and a comparison of the observed responses with the predicted ones setting up and all these batches were found close to the predicted values and show validity of optimized data.

Conclusion: Hence QbD approach is quite efficient to get optimized drug delivery systems og L-Arginine without doing exhaustive study.

Key words: L-Arginine, Gelatin, Central Composite Design, microspheres, Characterization of microspheres.
verimi ve ilaç salınımı ile yapıldı. Jelatin ve etil selüloz konsantrasyonunun etkisi, yüzde verimi, mikro küre boyutu ve ilaç tutma verimi gibi bağımlı tepki üzerinde belirlenmiştir.

**Bulgular:** Yanıt merkezi eğrisi 32 merkezi kompozit dizayn kullanılarak elde edildi ve sırasıyla yüzde verim, mikro küre boyutu ve ilaç tutma verimi 89.98, 333.32 mm ve% 82.61 olarak optimize edilmiş toplu elde edildi. Optimize edilmiş partinin geçerliliği, bağımsız yanıtın optimize edilmiş değerleri ile dört farklı parti formül e edildi ve gözlemlenen yanıtların öngörülen değerleri karşılaştırılması ve tüm bu partilerin öngörülen değerlerin yakınında bulunduğu ve düzgünleştirilmiş verilerin geçerliliğini gösterdi bulundu.

**Sonuç:** Sonuç olarak, QbD yaklaşımı, aşırı etkili çalışma yapmadan L-Arginin'in optimize edilmiş ilaç verme sistemlerini elde etmek için oldukça etkilidir.

**Anahtar kelimeler:** L-Arginin, Jelatin, Merkezi Kompozit Tasarımı, Mikrosferler, Mikrosferlerin Karakterizasyonu.

**Introduction**

Quality by Design (QbD) is helpful tool for systemic development of drug formulations based on sound scientific principles, hence it refers to the successful achievement of predictable quality with desired predetermined specification and without doing exhaustive conventional study.1 QbD paradigm of drug regulation necessitates very well understanding of the product to overcome future product failures.2 Design of Experiment (DoE) and RSM helps in finding the individual as well as combined effect of variables on product.3,4

Oral controlled release formulations are developed to improve the problems associated with oral conventional dosage forms like they can reduce side effects, improve the therapeutic efficacy by delayed/prolonged drug release so that frequency of drug administration can be reduce. Thus assuring better patient compliance.5,6 Various technique have been developed for controlled release formulations; which utilizes the cros-linking ability of polyelectrolytes in the presence of counter ions to form multiparticulate system. These delivery systems are spherical crosslinked hydrophilic polymeric system which upon gelation and swelling in simulated biological fluids releases drug in controlled manner. These developed microsphere are can be loaded with high amount of drug as compared to the conventional delivery system.7,8

Arginine an ergogenic (i.e., performance enhancing) supplement, most notably in the “nitric oxide” (NO) class of supplements is a semi-essential amino acid involved in multiple areas of human physiology and metabolism. NO produced from it improves outcomes in various diseases.9 L-arginine is readily available over the counter and is popular as a nutritional
supplement to increase muscle mass. More recently, L-arginine has been tested as a potential therapeutic in numerous acute and chronic disease states, including sickle cell chest crisis, pulmonary artery hypertension, coronary heart disease, pre-eclampsia and myocardial infarction, because of its bronchodilator and vasodilator actions.\textsuperscript{10-11}

**Materials**

L-Arginine was obtained from CDH Laboratory Chemicals, Sodium Alginate (low viscosity grade, 250 cp of 2% solution at 25°C) from Loba cheime Pvt Ltd (Mumbai). Gelatin, Ethyl Cellulose and Span 80 were purchased from Thermo fisher scientific India Pvt. Ltd. (Mumbai). Glutaraldehyde and Light liquid paraffin were procured from Loba chemical, Mumbai. All other chemicals used in the study are of analytical grade. HPLC grade water, methanol and potassium dihydrogen orthophosphate purchased from Qualigens fine chemicals (Gujrat).

**Methods**

**Preparation of microspheres**

Controlled released microspheres of L-arginine were prepared by performing cross linking of gelatin using glutaraldehyde. The required amount of gelatin was taken in a beaker; to this 8ml of distilled water was added and this mixture was heated at 40°C temperature for 3-4 min to get uniform polymer mixture. Different concentration of ethyl cellulose (EC) was added as shown in table-1. Then the specified amount of drug was dispersed thoroughly to the polymer solution. A mixture of Light liquid paraffin (200mL) and span 80 (0.1mL) was prepared. The mixture was maintained at 4°C with ice bath and stirred at 200 rpm and to this mixture previously prepared polymeric drug solution was added through a syringe with 22 gauge needle. After some time glutaraldehyde (2mL) was added drop wise to it with continuous stirring for 2 h. Microspheres were filtered, washed by iso-propyl-alcohol to remove liquid paraffin and dry at room temperature. Then dried microspheres were collected, weighed and stored.\textsuperscript{12-13}

**3\textsuperscript{2} Central Composite Design**

A 3\textsuperscript{2} CCD was adopted for optimization study. Two independent variables investigated were functional excipients such as concentration of gelatin (X) and EC (Y). The impact responses of these independent variables were investigated on the dependent responses such as percentage Yield, Microspheres Size (MS) and DEE. The experimental points used according to the design shown in Table 1.
Polynomial equations were generated and used to express the function of independent variables. Common polynomial equation to observe the effect of independent variable can be expressed as

\[ Y_1 = b_0 + b_1 X_1 + b_2 X_2 + b_3 X_1X_2 + b_4 X_{12} + b_5 X_{22} + b_6 X_1X_{22} + b_7X_{12}X_2 \quad \text{eqn.}(1) \]

Where \( Y_1 \) is the dependent variable, \( b_0 \) is the arithmetic mean response of the thirteen runs. The main independent variables, that is, effects \( X_1 \) and \( X_2 \) represent the average result of changing one factor at a time from its lower values to its higher values. \( 3^2 \) CCD is most efficient tool in estimating the influence of individual variables (main effects) and their interactions using minimum experimentation. In the present research, \( 3^2 \) CCD was considered to be best as the values of the response surfaces were not known from the previous findings. Thus, this design was selected for optimization of formulated microspheres.

**Evaluation of prepared microspheres**

**Characterization of microspheres**

FTIR spectra was obtained by Jasco FTIR 6100 type A, Japan spectrometer, sample was prepared in KBr disks, and spectra was recorded over the wavenumber 4000-400 cm\(^{-1}\). All three spectra were completely analyzed.\(^{14}\)

**Percentage yield**

Microspheres dried at room temperature were weighed and the Percentage yield of microspheres was calculated using formula.\(^{14}\)

\[
\% \text{ yield} = \frac{\text{Amount of sphere prepared experimentally}}{\text{Theoretical amount of microspheres (mg)}} \times 100 \quad \text{eqn.}(2)
\]

**Morphological analysis**

Scanning electron microscope (Zeiss, Supra 40, India) was used to characterize surface topography of the microspheres. The microspheres were fixed on a brass support with a thin adhesive tape and the samples were coated with thin layer gold under vacuum to render them electrically conductive (approximately 3000 Å). The surface picture was taken screened taken at 15kV and 20kV for the drug-loaded microsphere.

**Particle Size determination of microspheres**

Particle size analysis was done by sieving method. Microspheres were separated out in different size fractions by passing them through a set of sieves for 5 minutes. This set of sieves included standard sieves having nominal mesh apertures of 1.0 mm, 0.71 mm and 0.5 mm (sieve no.16, 22 and 30 respectively). The particle size distributions of the beads were determined and mean particle sizes of beads were calculated using following formula.
Mean particle size = \( \frac{\sum (\text{Mean particle size of the fraction} \times \text{weight})}{\sum \text{Weight fraction}} \)  

\text{eqn. (3)}

**Swelling index**

Gelatin microspheres were kept in double distilled water for swelling for 1h to reach maximum swelling. Volumetric measurements were done by determining the increase in volume in the swelling medium at specific time intervals. The swelling index was calculated as\(^{15}\):

\[
\text{Swelling Index} = \frac{\text{Volume of swollen particles}}{\text{Volume of dry particles}}
\]

\text{eqn. (4)}

**Drug Entrapment Efficiency**

Accurately weighed drug-loaded microspheres equivalent to 100 mg of L-Arginine were added to 0.1 N HCl and kept for shaking on mechanical shaker for 24 h. Then the solution was filtered and the drug content was estimated spectrophotometrically using HPLC (Younglin, ACME-9000, China). The drug entrapment efficiency was determined using following formula\(^{16}\):

\[
\text{Drug Entrapment Efficiency} = \frac{\text{Actual drug content}}{\text{Theoretical drug content}} \times 100
\]

\text{eqn. (5)}

**in-vitro Drug Release**

In vitro release studies were carried out on L-Arginine loaded microspheres using USP XXIV dissolution test apparatus-I (Electrolab, TDT-06T, Maharashtra, India). Weighed quantity of microcarriers equivalent to 100 mg of L-Arginine were introduced into a dissolution basket and the basket was placed in 900 mL of phosphate buffer solution (pH = 7.4 for 8 h) at 37 ± 0.5°C (Ph. US 24th edn) and 50 rpm.\(^{17}\) Aliquots of 5 mL solution were withdrawn at specific time intervals and replaced with fresh dissolution medium. The withdrawn samples were analysed for drug content at HPLC (YOUNGLIN, ACME-9000, China) having UV detector. The samples were studied at 210 nm to obtain the retention time 2.2-2.4 m and AUC.\(^{18}\) The results of in vitro release data were fitted into various release equations and kinetic models.\(^{19-21}\)

**Optimization and Data validation**

Concentrations of gelatin were selected as 1000, 900 and 800 mg; whereas for EC were 50, 100 and 150 mg. Thirteen formulation was developed by selecting nine possible combination among which centre point was repeated four times and mean value was taken for further study. The dependent responses were analyzed using Design Expert® 8.0.7.1 (trial version). The models were tested for significance and optimized batch was selected with desired values.
of dependent responses. Three formulations (VCB1 to VCB4) along with optimized batch were developed and validated by response surface methodology. The observed and predicted responses were critically compared. Linear correlation plots were constructed for the chosen check-point formulations. The residual graphs between predicted and observed responses were also constructed separately and the percent prediction error (% bias) was calculated with respect to the observed responses. Optimized batch was validated taking total three formulations selected as check-points.

Result
FTIR profile of formulated microspheres was done to identify the drug polymer interaction. Hence formulated microspheres were subjected to IR analysis to evaluate possible interaction between drug and polymer. Infra red curve of pure drug and formulated microspheres shows similar peak (Table-2) which confirms that there is no interaction between drug and polymer.

Surface topography
Scanning electron microscopy was used to investigate the surface topography of prepared microspheres and is shown in Figure. 1.

The percentage yield and Mean particle size of the formulation were depicted in table-3.

Drug Entrapment Efficiency
Drug Entrapment efficiency is an important variable used to assess the drug loading capacity of Microspheres and their drug release profile. DEE depends upon various parameters such as process used for preparation, physicochemical properties of the drug and various formulation variables (Table-3).

In vitro release behaviour of drug:
In vitro drug release behaviour of formulated glutaraldehyde cross-linked gelatin microspheres is shown in Figure-3. All batches were studied for their drug release profile for up to 8 h it was observed that in all formulated system the rate of release was varies due to use if different concentrations of dependent variables. It is clear that as amount of EC increases from 50-150 mg rate of drug release decreased which indicate hydrophobic nature of EC in the formulation hence increasing amount of Ethyl cellulose lead to retardation in drug release. Cross linking property of glutaraldehyde leads to formation of a rigid hydro gel to restrict the leaching thereby decrease the drug dissolution (Table 4 & Figure 3 ).

Data analysis and optimization
Drug release mechanism was investigated by fitting to models representing zero-order, first order, Higuchi’s square root of time model and Korsmeyer-Peppas model. Results of ANOVA for Response Surface Quadratic Model for various dependent parameters are:

% yield = +80.47+5.45A+2.73B+0.97AB+0.51A^2+0.32B^2-0.48A^2B-0.037AB^2 \hspace{1cm} \text{eqn. (7)}

MS = +317.64+11.15A+2.77B-0.62AB+0.29A^2+0.82B^2+1.10A^2B+0.31AB^2 \hspace{1cm} \text{eqn. (8)}

DEE= +76.38-0.075A+2.39B-0.67AB+1.79A^2+0.082B^2+0.70A^2B-0.48AB^2 \hspace{1cm} \text{eqn. (9)}

Where A indicates concentration of gelatin while B represents to concentration of EC.

**Validation of the statistical model**

Validation of optimized batch was done by formulating four different batches using overlay plot (figure 5) by utilizing the optimum value as founds by statistical tool i.e. by considering the optimum value as found (Table 5) and a comparative study was done between predicted value and observed values to determine the prediction error (Figure 6).

**Discussion**

FT-IR spectra (figure 2) of pure L-arginine and formulated microspheres of L-arginine shows the identical peaks as that of standard L-arginine which proves that excipients incorporated in formulated microspheres do not interact with L-arginine and all ingredients of beads are compatible with each other.

Scanning electron microscopy of microspheres of L- Arginine shows well-rounded spheres with rough surface because of sudden cross linking of gelatin with glutaradehyde. The particle size of the formulations was found to be between 320- 351.11μm. It was observed that the mean particle size of formulated microspheres were decreased with respect to the increased the amount of ethyl cellulose in the formulation.

Results for drug entrapment efficiency indicates that as the concentration of EC increases the DEE increase which is due practically insoluble nature of hydrophobic polymer i.e. EC. DEE was increased as the amount of EC was increased in the formulation because of practically insoluble nature of EC in water.

In vitro drug release study of microspheres of L- Arginine was carried out in 900 mL of phosphate buffer solution (pH = 7.4 for 8 h) at 37 ± 0.5°C. In the fasted state gel microcarriers exhibited a biphasic release profile as an initial rapid drug release phase due to burst which are loosely into or just beneath the surface of microspheres. followed by a
slower, gradually decreasing drug release phase after 1 hour extending up to 8 hours (Table 4 and Figure 3).

Drug release mechanism was investigated for number of models i.e. zero-order, first order, Higuchi’s square root of time model and Korsmeyer-Peppas model. zero-order, first order, Higuchi’s square root of time and Korsmeyer-Peppas model gave $R^2$ value 0.9101 to 0.9473, 0.9832- 0.9889, 0.9901- 0.9992 and 0.963-0.991 respectively, showing Fickian diffusion involving a combination of swelling, diffusion and/or erosion of matrices. Various response surface plots were also drawn to analyse impact of independent variables on dependent variables as discussed earlier.

Optimizations of formulated microspheres were done by using $3^2$ CCD. The outcomes for response parameters, that is, %Yield, MS and DEE were subjected to regression analysis and statistical models were found to be significant. Observed dependent responses i.e. %Yield, MS and DEE shows fair relation between the dependent and independent variables. Percentage yield for formulated batches were found in the range of 74.65– 86.36% while particle size was found in range of 302.34-.333.32 mm. Drug entrapment efficiency of all the formulation was found to be between 73.59 ± 1.744 to 82.61 ±0.700 Figure 4(a) shows that values of % Yield, increases with increase in concentration of gelatin and also increases with increasing EC concentration. Maximum % Yield is observed at the highest levels of Gelatin and EC.

Figure 4(b) shows a nearly linear ascending pattern for MS, as the content of gelatin increased, this MS increase slowly with increasing gelatin. Value of MS achieves to its maximum value at the highest levels of gelatin and EC. Nonlinear pattern of contour lines indicates significant impact on gelatin and EC.

Figure 4(c) shows that the DEE increases almost linearly with increase in concentration of gelatin whereas it decreases very slowly and then increases with increase in EC concentration. Maximum value of DEE was observed at the highest gelatin and EC concentration.

Validation of optimized batch was done by setting up a comparison of the observed responses with the predicted ones (Table 5), the prediction error varied between -0.024 to 0.048, -0.990 to 0.090 and -0.013 to 0.159 for %Yield, MS and DEE respectively. The linear correlation plots drawn between the predicted and observed responses, forcing the line through the origin, demonstrated high values of R (0.996 to 0.999, Figure 6), indicating excellent goodness of fit ($p < 0.005$). The corresponding residual plots show nearly uniform and random scatter around the mean values of response variables.
Conclusion:
Spherical amino acid loaded gelatin microspheres were prepared to achieve sustain release by cross-linking technique. Successful establishment of L-Arginine stacked repressible release matrix delivery system was done and escalation was achieved by using QbD applying $3^2$ central composite design. Hence it can be concluded that QbD is a powerful tool for present research that helps in developing desired formulation without wastage of time as well as man, money and material.

Reference