Preparation, characterization and scale-up of sesamol loaded solid lipid nanoparticles

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Abstract

Sesamol loaded solid lipid nanoparticles (S-SLNs) were prepared with the aim of minimizing its distribution to tissues and achieving its targeting to the brain. Three scale-up batches (100×1 L) of S-SLNs were prepared using a microemulsification technique and all parameters were statistically compared with the small batch (1×10 mL). S-SLNs with a particle size of less than 106 nm with a spherical shape (transmission electron microscopy) were successfully prepared with a total drug content and entrapment efficiency of 94.26±2.71% and 72.57±5.20%, respectively. Differential scanning calorimetry and infrared spectroscopy confirmed the formation of lipidic nanoparticles while powder X-ray diffraction revealed their amorphous profile. S-SLNs were found to be stable for three months at 5±3°C in accordance with International Conference on Harmonisation guidelines. The SLN preparation process was successfully scaled-up to a 100x batch on a laboratory scale. The procedure was easy to perform and allowed reproducible SLN dispersions to be obtained.

Introduction

Sesamol is the main antioxidant present in sesame oil. It is generated from sesaminol by roasting sesame seed or bleaching sesame oil. The antioxidant potential of sesamol has been established in vitro and its anti-inflammatory, chemoprotective, antiaging, cardioprotective and hepatoprotective effects are also well documented.

The physicochemical nature of sesamol (log P 1.29; solubility 38.8 mg/mL) substantially enhances its distribution to the tissues, thus minimizing its delivery to the brain. This led us to look for an alternative controlled release drug carrier system such that the brain could be efficiently targeted.

Solid lipid nanoparticles (SLNs) consist of spherical solid lipid particles in the nanometer range dispersed in a solution of surfactant which facilitates their passage across the biological membrane. Although there are numerous methods for the preparation of SLNs, only a few of them can be easily scaled-up.

The high pressure homogenization technique, which has been used since the 1950s for the production of parenteral emulsions, is the most common technique, and is used to produce large volumes of SLN dispersions. However, this manufacturing process requires large energy resources. Furthermore, the process is not suitable for the production of water soluble drugs that may partition towards the water phase as a result of the homogenization pressure, resulting in low entrapment efficiencies.

Still another efficient and robust method of producing SLNs is by microemulsification. This involves SLNs being prepared by the dispersion of warm oil-in-water (o/w) microemulsion in cold water (0-4°C); solid lipids with high melting points forming the internal phase of the microemulsions. Although scaling-up of this method for parenterals was reported by Marengo et al. in 2000, they scaled-up the process for 1 mL of microemulsion to only 100 mL using especially designed apparatus. The warm liquid microemulsion was sterilized through a 0.22 mm membrane filter; the nanodroplet size of warm o/w microemulsion is lower than 100 nm. After passing through the filter, the warm microemulsion was made to flow through the interchangeable needle and drop directly into a stirred cold aqueous medium. Small batches of 5 mL microemulsion poured into 50 mL of cold water (1:10 dilution) were being produced in our laboratory before this. The pharmacodynamic study of the prepared SLNs in an ovariecotomized rat model proved our intention to deliver significant amount of sesamol to brain.

However, in order to test the commercial viability of the process, we planned to develop the scaled-up batches of 1 liter (L) or more and characterize the SLN formulation that develops according to various in vitro parameters, including particle size, percentage entrapment efficiency (%EE), transmission electron microscopy (TEM), infrared spectroscopy (IR), differential scanning calorimetry (DSC), powder X-ray diffraction studies (PXRD) and in vitro release. We aimed to produce three scaled-up batches (1 L each) of sesamol SLNs using a microemulsification method and appropriately characterized them. Further objective of the study was to obtain a concentrated dispersion of nanoparticles by effecting only a 1:1 dilution of the hot microemulsion with cold water instead of a commonly 1:10 or 1:50 dilution. Production of a highly diluted SLN dispersion is one of the major limitations of the microemulsification method. The method requires subsequent steps such as diafiltration or lyophilization in order to obtain a stable concentrated product, followed by reconstitution for compliant usage. Both lyophilization and reconstitution can result in aggregation of particles to a micron size. So the other very important consideration of the study was to try and develop a one-step, concentrated and ready to use dosage form without any further treatment.

Materials and Methods

Materials

Sesamol was obtained from Jubilant Life Sciences (Noida, Uttar Pradesh, India); soy lecithin (Hi Media, Mumbai, India), Tween 80 (S.D. Fine Chemicals Ltd., Mumbai, India), and Compritol® 888 ATO (glyceryl behenate, a gift sample from Gattefosse, Paramus, NJ, USA) were also used in the study. All other chemicals and reagents were of analytical grade and were used without further purification.

Preparation of scaled-up batches of sesamol loaded solid lipid nanoparticles

Small batches (1×) of sesamol loaded solid lipid nanoparticles (S-SLNs) were produced using 10 mL glass beakers to maintain the lipidic and aqueous phases at temperature of 80-85°C. Suitable quantities were taken to produce a 5 mL of microemulsion which was poured into equal volume of ice-cold (0-4°C) water under magnetic stirring for the formation of solid lipid nanoparticulate dispersions. For production of a 100x batch of S-SLNs, 500 mL of microemulsion was produced. This was poured into an equivalent amount of ice-cold water (300 mL) to provide a concentrated dispersion. All the ingredients were weighed proportionally to produce a 100x batch. The process involved placing polysorbate 80 (250 mL) soy lecithin (80 mL), and water (180 mL)
Characterization of sesamol loaded solid lipid nanoparticles

Particle size analysis
The mean diameter of SLNs in the dispersion (with appropriate dilutions with triple distilled water) was determined using photon correlation spectroscopy (Zeta sizer 2000, Malvern Instruments Ltd., Malvern, UK).

Transmission electron microscopy
SLN morphology was examined using an electronic transmission microscope (Hitachi H-100; Hitachi Kokusai Electric Inc., Tokyo, Japan).

Differential scanning calorimetry
DSC was performed with a Perkin-Elmer differential calorimeter. DSC is a tool to investigate the melting and crystalline behavior of materials like SLNs. The breakdown or fusion of the crystal lattice by heating or cooling the sample provides information about the internal polymorphism, crystal ordering or glass transition processes. It uses the fact that different lipid modifications possess different melting points and enthalpies. The thermal analysis of the pure drug, lipid and SLNs were carried out to look for any significant changes in the pattern of the peaks. Samples were placed in a conventional aluminum pan and heated from 10°C to 250°C at a scan speed (5000 rpm) for exactly 2 h. SLNs are formed in the aqueous medium by crystallization of high melting point lipid droplets that represent the oil phase of the microemulsion. The prepared SLNs were stored in a refrigerator until further analysis. Three scaled-up batches were produced to check reproducibility of the selected method. All three were characterized using suitable parameters to ensure complete in vitro characterization and the results were also compared for uniformity of the parameters with the 1x laboratory scale batch.

The peaks obtained for the free drug and lipid, and their physical mixture was compared with the lyophilized SLN formulation to look for any significant changes.

Powder X-ray diffraction
The crystalline/amorphous nature of formulated nanoparticles was confirmed by X-ray diffraction measurements carried out with an X-ray diffractometer (XPERT-PRO, PANalytical, Almelo, The Netherlands). PXRD studies were performed by exposing the samples to CuKα radiation (45 kV, 40 mA) and scanning from 5° to 50°, 20 at a step size of 0.017° and scan step time of 25 s. Samples used for PXRD analysis were the same as those used for DSC. The instrument measures interlayer spacing d which is calculated from the scattering angle θ, using Bragg’s equation nλ = 2d sin θ where λ is the wavelength of the incident X-ray beam and n is the order of the interference. Obtained XRD patterns were compared for characteristic drug peak intensity.

Total drug content
Total drug content (TDC) was estimated by spectrophotometry at λ

\[ \text{λ}_{\text{max}} = 294 \text{ nm} \]

by disrupting 1 mL of the SLN dispersion using an appropriate volume of chloroform: methanol (1:1).

Entrapment efficiency
A dialysis bag with a cut off of 12 KDa (Hi Media) was used to estimate the EE. We poured 1 mL of S-SLN dispersion (3.72 mg/mL) in the dialysis bag, both ends of which were tied tight to prevent any leakage. The bag was dipped in 100 mL of water stirred magnetically at 150 rpm. Dialysate was withdrawn after 15 min and analyzed by spectrophotometry; time was optimized using an equivalent amount of free sesamol to be released in water; this was performed assuming no amount of sesamol is entrapped within the SLNs.

\[ \% \text{EE} = \frac{\text{Total drug} - \text{Drug in supernatant}}{\text{Total drug content}} \times 100 \]

Absorbance value obtained for blank SLNs treated in a similar manner was used as the control value to compensate for any interference with the ingredients. All measurements were taken in triplicate for all the three batches produced. Amount of drug in the dispersion retained inside the dialysis bag gave a direct measure of the quantity of drug entrapped.

Values obtained for the amount of drug in the supernatant and that retained in the dispersion inside the dialysis bag were added together to confirm the mass balance.

In vitro drug release
The in vitro release studies were carried out by the dialysis membrane method for studying the release of sesamol from the developed SLNs. The receptor media used for the studies was made up of 100 mL of phosphate buffer, pH 7.4, pre-equilibrated at 37±0.5°C. Aqueous SLN dispersion (1 mL containing 3.72 mg of sesamol as per the drug content studies) was placed in the dialysis tubing which was then sealed at both ends and dipped into the receptor media, maintained at 37±0.5°C and stirred continuously at 150 rpm. Samples of 3 mL each were withdrawn from the receptor medium with replacement at various time intervals and analyzed by spectrophotometry at 294 nm. All measurements were taken in triplicate for the three scale-up batches and also for the 1x lab-scale batch. Results obtained were analyzed statistically to confirm the reproducibility of the large scale batches produced using the microemulsification technique.

Stability
S-SLNs were stored in vials at 5±3°C for three months and the samples were withdrawn at 0, 1 and 3 months, as per International Conference on Harmonisation (ICH) guidelines. The average size, total drug content and the entrapment efficiency were determined at each time point.

Results
Characterization of sesamol loaded solid lipid nanoparticles

Particle size
Average particle size of all the three 100x batches was found to be 106.6 nm or under (Figure 1) with the actual size being 95.35, 98.11 and 106.6 nm for batches I, II and III. PI values were 0.432, 0.482 and 0.303 for the three batches, respectively. Particle size and the PI obtained for the 1x batch were 122 nm and 0.255, respectively.

Transmission electron microscopy
SLNs were found to be spherical in shape by TEM (Figure 2). The size of the lipidic nanoparticles observed under TEM (40-90 nm) was close (though smaller) to the results obtained using photon correlation spectroscopy.

Differential scanning calorimetry
DSC is a thermoanalytical technique in which the difference in the amount of heat required to maintain the sample and reference at same temperature is measured as a function of temperature and time. The basic principle underlying this technique is that when the sample undergoes a physical transformation together in a beaker which was heated to the lipid melt temperature. Glycerol behenate (40 g) was melted separately at 82-85°C. Sesamol (1:10 with respect to lipid) was added to the aqueous phase, following which the hot aqueous emulsifier mix was dropped at once into the lipid melt under magnetic stirring to obtain a clear microemulsion. The hot microemulsion thus formed was poured slowly in a streamline into an equivalent amount of cold water (~2°C) under continuous mechanical stirring (5000 rpm) for exactly 2 h. SLNs are formed in the aqueous medium by crystallization of high melting point lipid droplets that represent the oil phase of the microemulsion. The prepared SLNs were stored in a refrigerator until further analysis. Three scaled-up batches were produced to check reproducibility of the selected method. All three were characterized using suitable parameters to ensure complete in vitro characterization and the results were also compared for uniformity of the parameters with the 1x laboratory scale batch.
(such as melting, desolvation), some amount of heat is required to flow through it, depending on whether the process is exothermic or endothermic, to maintain both reference and sample at the same temperature. DSC measures this heat flow into or from the sample when it is heated or cooled. These measurements provide qualitative and quantitative information about physicochemical changes (i.e., endothermic, exothermic processes or changes in heat capacity). Figure 3 shows DSC thermograms of pure sesamol, Compritol® 888 ATO and the formed SLNs. The DSC curve for pure sesamol showed a fusion endotherm at 65.43°C corresponding to the melting point of sesamol (60-65°C) while the physical mixture shows peaks at 71.13°C (corresponding to the lipid) and another peak at 162.28°C (corresponding to the degradation peak of sesamol). Pure Compritol® 888 ATO shows an endothermic peak at 73.06°C. It also showed a degradation peak at over 160°C. However, DSC of developed SLNs did not show a peak corresponding to sesamol, confirming the successful incorporation of the drug into the SLNs. A broad endotherm starting from 95.17°C and showing two distinct peaks at 99.97°C and 112.36°C was observed for the SLNs, although a small shoulder was observed near the lipid melting point of 73.06°C. Observation of broad peaks indicates the amorphous nature of the developed SLNs. DSC for the laboratory scale (1x) batch and the 100x batch were similar.

Infrared spectroscopy

The IR peaks obtained with the developed formulation of S-SLNs revealed an intermolecular stretching of the -OH group (3400-3200 cm⁻¹) of the drug when compared with the peak of pure sesamol (Figure 4). This is a direct indication of the formation of SLNs as the stretching could not be observed when the physical mixture of the same components were analyzed (Figure 4). Thus, the formation of SLNs could be confirmed by IR analysis.

Powder X-ray diffraction studies

Overlaid PXRD patterns of sesamol, Compritol® 888 ATO, lyophilized blank SLN (BSLN) and S-SLN are shown in Figure 5. The PXRD pattern of sesamol showed sharp peaks at 2θ scattered angles 7.12, 15.19, 18.08 and 18.360 which indicated its crystalline nature. However, no characteristic peaks for sesamol in lyophilized S-SLNs were observed, indicating the amorphous nature of the formed SLNs. PXRD pattern of Compritol® 888 ATO shows sharp peaks at 2θ scattered angles 21.16, 23.37, 23.52 and 35.760; indicating the crystalline state of Compritol® 888 ATO.

Total drug content and entrapment efficiency

TDC and EE of S-SLNs (100x batch) was...
estimated to be 94.26±2.71% and 72.57±5.20% (n=9), respectively, and there was no significant difference (P≤0.05) between the TDC and EE of the three batches produced and the small scale batch (1x) (TDC-91.28±3.28%; EE-67.28±3.86%). High (EE) values (~70%) indicate the efficiency of the method for preparation of sesamol loaded SLNs and over 90% TDC confirms drug losses during formulation were insignificant. A significant EE also shows the suitability of the components and their relative proportion in the formulated SLNs.

**In vitro drug release**

The drug release from S-SLNs at 37±0.5°C is shown in Figure 6. The release of sesamol from S-SLNs was fitted to a first order kinetics model and occurs by diffusion. The release was prolonged for up to 24 h with approximately 50% of the drug being released before 4 h (Figure 6). Almost 90% of the drug was released in less than 16 h which may be due to the water soluble nature of sesamol and hence its incorporation into the outer phospholipid layer of the lipidic nanoparticles. However, a prolonged release was observed in the later stage (up to 24 h) which may be attributed to the diffusion of the remaining drug from the lipidic core. Overall, the system followed a non-Fickian drug release. In case of the 1x batch, a slower release was obtained and a total release of 68% was achieved at the end of 24 h.

**Stability**

After three months of storage at 5±3°C, the S-SLNs were found to be stable according to ICH guidelines without any significant increase (P≤0.05) in particle size (Table 1). Change in EE (5.2%) and TDC (2.2%) at three months was also without significance (P≤0.05) with respect to the 0 time samples, indicating a stable formulation.

**Discussion**

The main aim of this research was to scale up the preparation process of SLNs so as to obtain a highly stable concentrated dispersion. The study also aimed to design a process which can be carried out very easily and using relatively unsophisticated equipment. The use of the simplest equipment possible would provide scope for its commercial launch even by those conventional companies who specialize in simple pharmaceutical products such as emulsions. The phytochemical industry is also ready to venture into the development of phytopharmaceutical products that involve a minimum investment in sophisticated equipment. A stable microemulsion was formed even when it was prepared in a 100-fold (300 mL)
batch volume when compared to the small 1x laboratory batch. The microemulsion formed needs to be stable for a sufficient period of time to allow it to be poured into cold water as slowly and smoothly as possible under mechanical stirring (5000 rpm). This process can now be completed in 30 min, with 5-10 min of pouring time and 15 min of continued mechanical stirring, even after the complete addition of microemulsion. In the small scale (1x) batch, we were following stirring for long periods of time (3-4 h) after transfer of the microemulsion in order to achieve a small particle size.

In our previous experiments, o/w microemulsion was dispersed using a 18 gauge needle attached suitably to a glass syringe as reported and recommended (even for scaled-up batches) by other workers. Even though the process always resulted in a small particle size (<200 nm), it is demanding and difficult to use a hot syringe as the emulsion constitutes a high melting lipid, so any cool surface results in congealing of the lipidic phase; neither is the process suitable to be scaled-up easily on a laboratory scale. To overcome the problem, we tried to pour the microemulsion directly into cold water. We could successfully produce SLNs with the required particle size and a low polydispersity index, and the process could be suitably scaled-up to obtain a concentrated (1:1 dilution) SLN dispersion (1 L), with an average particle size as small as 95 nm (PI of 0.3) for one of the batches. However, PI is not a very important issue for formulations which are administered orally.

SLNs are formed by rapid quenching of the warm o/w microemulsion and the temperature difference between the warm microemulsion and the cold dispersing water plays an important role in determining the size of SLN. A rapid crystallization of the oil droplets of the warm microemulsion during quenching favors the formation of small SLNs, avoiding the coalescence among the oil nanodroplets.

The aqueous system made up of sesamol in polysorbate 80 and soy lecithin, while Compritol® 888 ATO was chosen as the lipid component as it results in dispersions with small particle size. The main reason for the selection of Compritol® 888 ATO as lipidic phase is its unique composition of glycerol triglyceride (28-32%, glycerol dibehenate (52-54%) and glycerol monobehenate (12-18%). The most abundant fatty acid is behenic acid (>85%) but other fatty acids (C16-C20) are also present. Compritol® 888 ATO has an amphiphilic character due to the presence of partial acylglycerols. Its hydrophilic lipophilic balance (HLB) is approximately 2, having a melting point between 69°C and 74°C and a density value of 0.94 g/cm³. Compritol® 888 ATO has a peroxide value lower than 6 meq O₂/kg indicating high chemical stability. It is thought that a lipid containing a major portion of diglycerides would be better able to hold an amphiphilic drug and would thus result in more efficient entrapment. Sesamol shows a solubility of 38.8 mg/mL and a log P of 1.29, favoring the octanol phase, thus showing it to be an amphiphilic molecule. The results obtained with the scaled-up batches of S-SLNs were found to be reproducible. Also, there was no difference between the characterization parameters of the 1x and 100x batches. There was no significant difference (P≤0.05) between the particle size, drug content and entrapment efficiency either within the scaled-up batches or the 1x batch, indicative of the robustness of the method. However, a sig-

![Figure 5. Overlaid powder X-ray diffraction studies patterns of Compritol® 888 ATO, sesamol, blank solid lipid nanoparticles (BSLN) and sesamol loaded solid lipid nanoparticles (S-SLNs).](image)

![Figure 6. In vitro release profile of sesamol, from sesamol loaded solid lipid nanoparticles, by dialysis.](image)

<table>
<thead>
<tr>
<th>Time points</th>
<th>Av. particle size (nm)</th>
<th>Total drug content (%)</th>
<th>Entrapment efficiency (%)</th>
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<tr>
<td>0 time</td>
<td>106±19.60</td>
<td>94.26±2.71</td>
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<td>1 month</td>
<td>95±22.00</td>
<td>92.70±4.16</td>
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<td>3 month</td>
<td>109±15.60</td>
<td>92.30±1.58</td>
<td>68.30±2.15</td>
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Table 1. Stability study parameters during storage of sesamol loaded solid lipid nanoparticles at 5±3°C.
significant difference (P≤0.05) was observed for the cumulative amount released for the 1x batch which was only 68% up till 24 h and the 100x batch (>90% at 24 h). This difference may be due to the amount of surfactant left unused after the formulation of SLNs. Scaled-up batches may accumulate a higher percentage of surfactant both on the surface of the formed nanoparticles and/or in the surrounding aqueous phase, helping the drug to diffuse out more rapidly, especially the drug entrapped in the inner lipidic core to the outer aqueous phase. The high reproducibility of results with respect to the characterization of SLNs may be due to an optimized formula for producing the microemulsion. Final composition of the microemulsion resulted in its spontaneous formation with an average particle size of less than 40 nm. When poured into ice cold water under high speed stirring this was expected to result in solid particles in the nanometric range. Stability of the formulation for up to three months was in accordance with ICH guidelines. Several cases report that SLNs show an inadvertent increase in particle size when kept and stored under refrigerated conditions, due to the gelling of the lipid phase. We, however, did not observe any such gelling phenomenon, which may be due to the high viscosity imparted by the concentrated SLN dispersion. A high concentration of polysorbate 80>20% may also contribute to the high viscosity which may also help the nanoparticles to remain in a dispersed form without any aggregate formation.

Also, in other methods used for scaling-up solid lipid nanoparticles, especially high pressure homogenization, in general, a large number of formulation parameters have to be considered before reaching an optimal formulation. Based on the data obtained, scale-up using microemulsification proved to be relatively easy, mainly because it is a spontaneous process and once appropriate proportions of the constituents (surfactant, lipid and water) are obtained they will always result in a microemulsion. Furthermore, the final particle size of the formed SLNs is mainly monitored by the particle size of the microemulsion which will only form when the particle size is in the nano range.

Conclusions

In conclusion, the SLN preparation process was successfully scaled-up to a 100x batch on a laboratory scale, was easy to perform and allowed reproducible SLN dispersions to be obtained. Research is ongoing for the second step of the scale-up design to produce batches of over 20 L, and then subsequently 200 L.

References

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