Review Article

Development of Cuboidal Nanomedicine by Nanotechnology

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Abstract

When amphiphilic lipid systems are placed in aqueous environment that will formed self-assembled nanostructured dispersed particles which are expressed as “Cubosomes” whose size ranges from 10-500 nm in diameter and they appear like dots square shaped, slightly spherical in the form of bicontinuous cubic liquid crystalline phase. The discovery of cubosomes is unique story and spans in the field of food science, biological membranes, differential geometry and digestive process. Cubosomal drug delivery system is great potential in melanoma therapy owing to their potential advantage. Recently the research interesting is inducing on cubosomes due to their thermodynamically stable cubic structure, biodegrability of lipids, the ability of encapsulating hydrophobic, hydrophilic, amphiphilic substances is an excellent candidate for targeting and controlled release of drugs, proteins etc. The review article includes manufacturing techniques, characterisation, and applications of cuboidal drug delivery system.

Keywords: Cubosomes; Cubic phase; Bicontinuous; Biodegrability; Controlled release

Introduction

Cubic liquid crystals are physically looks transparent and isotropic phases that are stable in excess water and show a unique system for the production of pharmaceutical dosage forms. The liquid crystals of cubic phase are used in the controlled release of selected water and oil soluble molecules [1].

Cubosomes are bicontinuous cubic liquid crystalline materials are active ingredients because they give the unique structural ends to control release applications [2]. Cubosomes are vesicle derivatives [3], formed by the mixture of phospholipid (glycerol monooleate) and non-ionic surfactant (poloxamer 407- block copolymers of polyoxyethylene polyoxypropylene) in aqueous media by applying high energy dispersion such as sonication and homogenization, after the formation of cubosomes the dispersion is formulated into a product and then applied to a substrate of interest usually body tissue [4], due to the formation of vesicles the stability of system remains in the interested area (figure 1) [5].

Structurally cubosomes are lipid vesicles formed from amphiphilic building blocks [6], which mimics bio membranes or vesicular structure similar to liposomes that can be used for carrier potential of hydrophilic, lipophilic and amphiphilic drugs as compare to free drug directly to the particular site of action [7], thus allow drug targeting and the sustained or controlled release of conventional medicines [8]. Cubosomes can be administrated in many ways oral, Percutaneous, intravenous route [9].

Among the various colloid carriers or vesicular delivery system, liposome and cubosomes can encapsulate both hydrophilic and lipophilic drugs. The hydrophilic drug is encapsulated inside the vesicles whereas the lipophilic drug is partitioned between the hydrophilic domains. Liposomes are produced by the self-assembly of phospholipids in aqueous phase to form bilayer which may be spherical unilamellar or multilamellar vesicles. Liposome may undergo some problems like degradation by hydrolysis in aqueous solution, sedimentation and aggregation on storage and cannot sterilize for clinical use. Still, chemical instability like oxidation of phospholipids was not avoided. These pave the way to the discovery of non-ionic surfactant vesicles known as cubosomes [10].

In several cases the conventional chemotherapy is not effective in the treatment of intracellular infection due to limited drug permeation into cells this can be overcome by the use of cubosomal drug delivery system [11], and also cubosomes reduces toxicity of encapsulated agent [12], dose related side effects, decreasing dosing frequency [13]. Like ethosomes cubosomal drug is administrated in semisolid form (gel or cream), producing high patient compliance rather than other vesicular drug delivery system [14]. With respect
to liposomes cubosomes possesses a larger ratio between the bilayer area and the particle volume and a larger breaking resistance [15]. The actual motto of cubosomes is to control degradation of drugs and loss, prevention of harmful side effects and increase availability of drug at the disease site [16].

One of the reasons for preparing cubosomes is the high chemico-physical stability of surfactant than that of phospholipids which are used in the preparation of liposomes. Due to the presence of phospholipids are easily hydrolysed [17]. Manufacture of cubosomes on a large scale embodied difficulty because of their viscosity [18].

Cubosomes are useful over other drug delivery system due to their bioavailability improvement of poorly soluble drugs and enhances skin permeation, by this reduces the cost of therapy [19]. Surfactant used in cubosomes does not require special handling and storage condition [20].

Cubosomes have a transdermal effect of which has been reported by some researchers, this effect might be due to their structural organization similar to that of biomembranes [21], recently the structural organization of the stratum corneum can be investigates by the study of cryo-transmission electron microscopy. Although cubosome structure has been characterized by X-ray and NMR investigations [22].

As described above cubosomes are nanoparticles with size range of 10-500 nm in dimension, they look like dots square, slightly spherical shaped (figure 2). Each dot shows the presence of pore having aqueous cubic phases in lipid water system in X-ray scattering technique was first identified by Luzzati & Husson [23].

The cubic phase structure of cubosomes can be described by the concept of differential geometry and periodic minimal surfaces by analogy with soap films. Mathematically ‘Schwarz’ discovered three types (figure 3) of minimal surfaces in cubic phases based on their curvatures.

In the monoolein water system the D-surfaces formed at high water levels, the G-surface at lower levels and the P-surface is formed when caseins or amphiphilic block copolymer are added [24].

Overall, cubosomes have great potential in drug nanoformulations for melanoma therapy owing to their potential advantages, including high drug payloads due to high internal surface area and cubic crystalline structures, relatively simple preparation method, biodegradability of lipids, the ability of encapsulating hydrophobic, hydrophilic and amphiphilic substances, targeting and controlled release of bioactive agents like proteins and drugs [25].

The aim of this work is to study the performance of cubosomes as cutaneous as well as oral delivery systems for Non-Steroidal Anti-Inflammatory Drugs (NSAIDs) seem particularly interesting. The NSAID’s shows their anti-inflammatory activity by obstructing the enzymes Cyclooxygenase (COX1 & COX2), which have to be proven an effective corticosteroid alternative in the topical management of percutaneous inflammation [26]. Presently NSAIDs are widely used in treatment of allergic conjunctivitis, management of post-operative inflammation and control of pain after photo refractive keratectomy. Practically NSAIDs are irritating due to their acidic nature. By reducing the pH of their formulation also further increases their irritation potential [27].

The first part of the present study describes the production and characterization of NSAID’s containing Monooleine (MO) dispersions, whereas the second part deals with in-vitro from different formulations using excised human skin membranes are Stratum Corneum Epidermis (SCE) mounted into Franz cells and an in-vivo investigation to describe the release of NSAID’s after administration (table 1).

**Properties of cubosomes**

1. Cubosome dispersions have much lower viscosity.
2. Cubosomes are discrete, sub-micron, nanostructured particles of bicontinuous cubic liquid crystalline phase.
3. Cubosomes is perhaps the most intriguing [29].
4. Cubic liquid crystals are transparent and isotropic phases that are physically stable in excess water.
5. Due to small pore size of cubosomes are attractive for controlled release.
6. It’s having an ability of solubilizing hydrophobic, hydrophilic and amphiphilic molecules and its biodegradability by simple enzyme [30].

**Advantages of cubosomes**
Cuboidal systems have many advantages for drug delivery which are as following,

1. They have ability to encapsulate both hydrophilic and hydrophobic also amphiphilic drugs.
2. They have a sustained release drug delivery characteristics.
3. Cubosomes have biocompatibility and bioadhesivity properties.
4. Bicontinuous cubic liquid crystalline phase of Cubosomes even stable in excess water [31].
5. Cubosomes are excellent solubilizers, compared with conventional lipid or non-lipid carriers.
6. They show high drug carrier capacity for a range of sparingly water-soluble drugs.
7. These are an excellent vehicle to protect the sensitive drug from enzymatic degradation and in-vivo degradation, such as peptides and proteins.
8. The cuboidal system enhances the bioavailability range twenty to more than one hundred times of water-soluble peptides [32].

Disadvantages of cubosomes

1. Cubosomes may lead to low drug loading efficiency and drug leakage in preparation, preservation and transport in vivo, thus the major problem of their stability acts as a barrier and thus limiting their use [33].

Advantages of cubosomes (phospholipids based carrier system) in comparison to other delivery systems

1. These systems show enhanced permeation of drug through skin for Percutaneous and dermal delivery.
2. These are platform for the delivery of large and diverse group of drugs (peptides, protein molecules).
3. Their composition is safe and the components are approved for pharmaceutical and cosmetic use.
4. Low risk profile- the toxicoological profiles of the phospholipids are well documented in the scientific literature.
5. High market attractiveness for products with proprietary technology. Relatively simple to manufacture with no complicated technical investments required for production of Ethosomes.
6. The vesicular system is passive and non-invasive, it is available for immediate commercialization (table 2) [34].

Materials used for preparing cubosomes

NSAID’s as a drug, glyceryl monooleate (monoolein), Poloxamer 407, Glycerol, Carbopol 934P, Alcohol (ethanol), Stabilizing agent (Polyvinyl alcohol), Bidistilled water, etc.

Techniques used for production of Cubosomes

The following techniques are used to produce cuboidal and nanocuboidal drug delivery systems,

<table>
<thead>
<tr>
<th>Sl.No</th>
<th>Cubosomes</th>
<th>Liposomes</th>
</tr>
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<tbody>
<tr>
<td>1.</td>
<td>Cubosomes are formation of bicontinuous cubic liquid crystalline phase by hydrating mixture of monolein and poloxamer 407.</td>
<td>Liposomes are formations of vesicles by hydrating mixture of cholesterol and phospholipids.</td>
</tr>
<tr>
<td>2.</td>
<td>Are artificial, colloidal and spherical vesicles of 0.05-5.0 μm diameter.</td>
<td>Are appear like dots square shaped, slightly spherical of 10-500nm in diameter</td>
</tr>
<tr>
<td>3.</td>
<td>In cubosomes active chemical constituent molecules are anchored through chemical bonds to the polar head of the phospholipids.</td>
<td>In liposomes, the active principle is dissolved in the medium of the cavity or in the layers of the membrane. No chemical bonds are formed.</td>
</tr>
<tr>
<td>4.</td>
<td>In cubosomes, polymer and the individual drug compound form a1:1 or 2:1 complex depending on the substance.</td>
<td>In liposomes, hundreds and thousands of phosphatidylicholine molecules surround the water soluble molecule.</td>
</tr>
</tbody>
</table>
Top down technique
Bottom up technique
Heat treatment
Spray drying

**Top down technique approach**

The viscous bulk cubic phase is prepared by mixing lipids with stabilizers, then the resultant mixture is dispersed into aqueous solution by the input of high energy (such as High-Pressure Homogenization [HPH], sonication or shearing) to form Lyotropic Liquid Crystal (LLC) nanoparticles (figure 4). HPH is the mostly used technique in the preparation of LLC nanoparticles. The cubosomes prepared through top-down approach are always observed to coexist with vesicles (dispersed nanoparticles of lamellar liquid crystalline phase) or vesicle-like structures [35].

**Bottom up technique approach**

In the bottom-up approach the hydrotrope is dissolve in water-insoluble lipids to create liquid precursors and prevent the formation of liquid crystals at high concentration and needs less energy input (Almgren et al). Discuss the formation of cubosomes by dispersing inverse micellar phase droplets in water at 80 °C and allow them to slowly cool, gradually droplets get crystallizes to cubosomes (figure 5). The cubosomes are spontaneously formed by emulsification. This bottom-up approach cannot effectively avoid forming vesicles through cryo-TEM, many vesicles and vesicle-like structures were also observed to coexist with cubosomes [36].

**Heat treatment approach**

This technique is not an integrated process for the manufacture of cubosomes because it only promotes the transformation from non-cubic vesicles to well-ordered cubic particles comprising a homogenization and heat-treatment step as a result decrease in the small particle size fraction that corresponded to vesicles and form more cubic phases with narrow particle distribution and good colloidal stability (figure 6) [37].

**Spray drying approach**

Because of the less flexibility of liquid precursor for formation of cubosomes (Spicer et al) developed dry powder precursor for cubosomes preparation. They utilized spray drying technique for preparation of starch encapsulated monoolein precursor and dextran encapsulated monoolein precursor. High proportion of polymer (75% w/w for starch and 60%w/w for dextran) for encapsulation decreased the amount of loading of active materials so system was limited for potent medicament, vitamins, flavours, or scents (figure 7 and 8) [38].

**General method of cuboidal preparation**

Cubosomes are usually produced by combining monoolein and water at 40°C. The resultant cubic liquid crystalline gel is dispersed into particles via the application of mechanical or ultrasonic energy. High-pressure homogenizers are often employed to produce cubosomes. Finally, the cubosomes are stabilized against flocculation.

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**Figure 4:** Preparation of cubosomes by top down approach.

- Lipid
- Aqueous phase
- Hydro trope
- LLC nanoparticles

**Figure 5:** Preparation of cubosomes by bottom up approach.

- Lipid
- Hydro trope
- LLC nanoparticles

**Figure 6:** Preparation of cubosomes by bottom up approach.

- Monoolein + aqueous solution of starch
- High shear mixture
- Coarse dispersion of cubosomes
- Spray drying
- Rehydrated cubosomes

**Figure 7:** Flow chart for formation of dry powder cubosomes.
by polymer addition (figure 9) [39].

**Evaluation parameters of cubosomes**

The cuboidal dispersions can be evaluated by following parameters,

- Thermal Analysis
- Polarizing Light Microscopy
- Cryo-Transmission Electron Microscopy
- X-ray Diffraction Measurements
- Drug Content of Dispersions
- HPLC Procedure
- In Vivo Studies
- Tape Stripping
- Statistical Analysis
- In vitro drug release studies

**Thermal analysis**

In this study, DSC is used to evaluate the physical status of the drug within the cubosomes. Both ingredients of the cubosomes seem to melt together in temperature of around 37°C to 56°C which may results of plasticizing of Glycerol Monooleate (GMO). The thermal events related to the drug’s melting point are different from those of the native drug (no sharp drug melting peak at around 200°C). The thermal events observed between 200°C and 300°C may be related to glyceryl monooleate degradation process [40].

**Polarizing light microscopy**

Samples were viewed between crossed polarizers and a λ-sheet in a Zeiss III light microscope (Zeiss, Oberkochen, Germany) [41].

**Cryo-transmission electron microscopy**

A small amount of prepared sample at ambient conditions is placed on a pure thin bar 600-mesh TEM grid. The solution blotted with filter paper to form a thin film. Then the sample is vitrified by immersing into liquid ethane near its freezing point. This is transferred to a transmission electron microscope for imaging using a cryoholder with temperature maintains 175°C. Images are recorded digitally by a charge coupled device camera using an image processing system [42].

**X-ray diffraction measurements (XRD)**

The XRD is carried out using a Philips PW 1830 X-ray generator. Diffraction patterns were recorded by using an INEL Curved Position Sensitive 120 detector. Diffraction data are collected at 25°C controlling the temperature.

Little structural information was derived from X-ray diffraction patterns: once the symmetry of the lipid phases was found, the unit cell dimension was calculated by using Bragg’s law as follows:

\[
\frac{1}{a} = 2\sin \theta / \lambda \\
\text{Where } a = \left( h^2 + k^2 + l^2 \right)^{1/2}
\]

Where \( h = 2 \sin \theta / \lambda \) with 2 \( \theta \) the scattering angle and \( \lambda \) (0.154 nm) the wavelength (2, 16) [43].

**Drug content of dispersions**

The drug content of dispersions evaluate by diluting the filtered dispersion sample in methanol (1:9 v/v) and analysed for drug content by High Performance Liquid Chromatography (HPLC) with the respective procedure. For Sedimentation Field Flow Fractionation (SdFFF) and stability studies, the amount of drug detected by HPLC after filtration was taken as reference of the total amount of drug [44].

**HPLC procedure**

Samples were assayed by using a validated HPTLC densitometric method. Developed plates were stained using a solution containing mobile phase cupric sulfate (pentahydrate): phosphoric acid: water and quantified densitometrically using a UV light source set respected wavelength [45].

**In vivo studies**

In vivo experiments were performed on ten volunteers for evaluation of anti-inflammatory activity. In Vivo Anti-inflammatory
Activity Ultraviolet-B-induced skin erythema was monitored by using a reflectance visible spectrophotometer X-Rite model 968 (X-Rite Inc. Grandville, MI, USA), calibrated and controlled. The reflectance spectra were obtained over the required wave length and 2-standard observer. From the spectral data of erythema, the Erythema Index (EI) was calculated using Eq. (1) [46].

$$E.I=100\log 1/R_{560} + (\log 1/R_{540} + \log 1/R_{580}) - 2(\log 1/R_{500} + \log 1/R_{510}) - 1$$

Skin erythema is induced by UVB irradiation using a UVM-57 ultraviolet lamp (UVP, San Gabriel, CA, USA) whose specific parameters are reported elsewhere. An irradiation dose is preliminarily determined twice the value of Minimal Erythemal Dose (MED) throughout the study [47].

Tape stripping

In this parameter the 200 mg of each hydrogel formulations is applied on defined cutaneous sites of forearm (three sites of application for each formulation induplicate). Spread the preparations uniformly by solid glass rod at the site and are then kept for 6h. After this period, the residual formulations are removed by wiping with cotton balls. Twenty individual 2 cm squares of adhesive tape are utilized on the application sites. Weigh each tape removed with stratum corneum cells then determination of stratum corneum removed by difference of weights; quantify the drug content in the tapes by some analytical technique [48].

Statistical analysis

This statistical analysis of in vitro data was performed using Student’s t test. Statistical differences of in vivo data are determined using repeated-measures analysis of variance (ANOVA) followed by the Bonferroni-YDunn post hoc pairwise comparison procedure. A probability of p<0.05 is considered significant in this study [49].

In vitro drug release studies

In vitro release of drug from cubosomes is evaluated using a dynamic dialysis method. The samples of various formulations were placed in dialysis bags (cellulose membrane), then immersed in 500 mL of simulated tear at 37±1 °C. Maintained the required paddle rotation speed. At predetermined time intervals, a 5 mL sample was withdrawn and immediately replaced with an equal volume of tear. Then the concentration of released drug is measured [50].

Stability studies on storage

The monoolein dispersions stored at room temperature leads to change with time. After preparation the dispersions are clear and transparent which will turn turbid with storage time. A number of macroscopically visible particles with sizes in the micrometer to millimeter range were formed during storage within days or weeks after preparation making these systems unsuitable for intravenous drug delivery. At refrigerator temperature the storage of homogenized dispersions results in the formation of white, semi-solid, ointment-like gels. This phenomenon is correlated with the crystallization of the colloidally dispersed monoolein. This type of behavior is observed in monoolein/water bulk systems due to high tendency for super cooling at sub-ambient temperatures. But the crystallization in the ternary monoolein/poloxamer/water bulk system could not be observed due to addition of poloxamer which modifies the crystallization behaviour of monoolein [51].

Applications of Cuboidal drug delivery system

1. Cubosomes are widely used in melanoma (cancer) therapy [52].
2. The monoglyceride based cubosome dispersion can be used for topically, such as for percutaneous or mucosal applications.
3. Due to microbiocidal properties of monoglycerides, can be used to design intravaginal treatment of sexually transmitted diseases caused by viruses (e.g. HSV, HIV) or by bacteria (eg. Chlamydia trachomatis and neisseria genorttiae).
4. The cubosomal technology is used to develop a synthetic vernix (complex mixture of lipid (fats), proteins and water) the cheesy white substance that coats infants in late gestation to help premature infants who are born without it. It is formed late in gestation and has an integral role in normal skin development.
5. Cubosome particles are used as oil water emulsion stabilizers and pollutant absorbents in cosmetics.
6. More recent use is as skin care, hair care, cosmetics and antiperspirants [53].

Future perspective

As discussed in this article, cubosomes contains drugs can be delivered to a patient by transdermal, oral and intravenously. With the sequence of human genome, biotechnology companies are developing a peptide and protein based drugs. It is expected that in the next 10 to 20 years, cubosomes containing protein and peptide based drugs will constitute more than half of the new drugs introduced into the market and more than 80% of these protein drugs will be antibodies due to control release activity. These biopharmaceuticals (proteins, peptides, carbohydrates, oligo-nucleotides, and nucleic acids in the form of DNA) present drug delivery challenges because these are often large molecules that degrade rapidly in the blood stream. Moreover, they have a limited ability to cross cell membranes and generally cannot be delivered orally. Such molecules will be much more difficult to deliver via conventional routes and injections may be the only means of delivery (at least as of today). The routes of administration will be dictated by the drug, disease state, and desired site of action. Some sites are easy to reach such as the nose, the mouth, and the vagina. Others sites are more challenging to access, such as the brain.

Conclusion

A simple process, based on dispersion and homogenization of monooleine and poloxamer 407 in water leads to the formation of a complex heterogeneous system, suitable for the delivery of lipophilic, hydrophilic drugs through the skin and use in treatments of hair, and other body tissue. The data here reported indicate a prolonged anti-inflammatory activity exerted by NSAID’s. The stratum corneum
like structure attributed to cubosomes by other authors suggests the hypothesis of a cubosome depot effect on the epidermis. Further specialized studies are required to confirm this fascinating hypothesis and to better investigate the role of vesicles and cubosomes in controlling the release of the drug.

References


