Transdermal drug delivery is an attractive route for the systemic delivery of actives since its first commercialization in 1979 [1]. Topical delivery via the skin is a convenient, painless, and non-invasive approach that enables self-administration for patients [2, 3]. Many drugs have been delivered via the transdermal route and have become successful at clinical setup [4, 5]. It is well established that the outermost layer of the skin poses a major obstacle in permeating the drug across the skin layer. Subsequently, the drug employed should possess certain physicochemical properties of low molecular weight and sufficient hydrophilic-lipophilic balance to successfully penetrate the skin [6]. Non-steroidal anti-inflammatory (NSAIDs) drugs possess favorable physicochemical properties and have been explored for the transdermal route [1, 7]. Among NSAIDs, Ibuprofen is the most commonly used and most prominently prescribed drug to alleviate moderate pain, reduce swelling, control pyrexia, and treat arthritis at higher doses [8, 9]. However, like other NSAIDs, Ibuprofen carries a risk of gastric ulcers and sometimes bleeding when taken orally.

Over the years, a variety of formulations have been explored to achieve transdermal delivery of a variety of therapeutic agents such as vesicle-based formulations, nanoparticles, microemulsions, and organogels and as so on [10]. Amongst them, organogels (OG) have attracted a great deal of attention due to their relevance in numerous applications such as transdermal drug delivery, separation science, templates, sensors, etc. They are thermoreversible bicontinuous systems consisting of oily surfactant solutions in which water is micro-emulsified, which on cooling develop a self-assembled interconnected network, mostly fibrillar or tubular, that stop macroscopic flow due to surface tension and capillary forces [11, 12]. The use of organogels has been proposed for the transdermal delivery of various drugs [13-16]. In this view, three major types of OGs comprising gelatine [17, 18], lecithin [19-21] and sorbitan esters [22, 23] have been studied. The OG formulations of our interest are based on non-ionic sorbitan ester surfactants. These belong to a class of surfactants that form lamellar, toroidal, tubular aggregates, and star-shaped clusters in OGs, which were evaluated using optical microscopy, freeze-fracture electron microscopy, and X-ray diffraction techniques. The drug release characteristics from sorbitan ester OGs are also reported [24].

Previously, Upadhyay et al. have evaluated phase behavior, rheological, and drug release characteristics of sorbitan ester OG to optimize the physical properties of the final product because of the equipment like planetary mixers for semisolid processing impart high shear that affects gelation [23]. In this work, we aim to formulate and characterize sorbitan ester OG consisting of Ibuprofen. First, we report the phase diagram for the OG formation at different ratios of hydrophobic/hydrophilic sorbitan ester surfactants consisting of drugs. Then, we investigated rheological characteristics and fractal dimension. Nonetheless, research on topical bases for the improvement in drug permeation is continued and in this paper, we showed that OG is a better alternative for prolongs the effect of Ibuprofen from conventional ibuprofen preparations.

**INTRODUCTION**

Transdermal drug delivery is an attractive route for the systemic delivery of actives since its first commercialization in 1979 [1]. Topical delivery via the skin is a convenient, painless, and non-invasive approach that enables self-administration for patients [2, 3]. Many drugs have been delivered via the transdermal route and have become successful at clinical setup [4, 5]. It is well established that the outermost layer of the skin poses a major obstacle in permeating the drug across the skin layer. Subsequently, the drug employed should possess certain physicochemical properties of low molecular weight and sufficient hydrophilic-lipophilic balance to successfully penetrate the skin [6]. Non-steroidal anti-inflammatory (NSAIDs) drugs possess favorable physicochemical properties and have been explored for the transdermal route [1, 7]. Among NSAIDs, Ibuprofen is the most commonly used and most prominently prescribed drug to alleviate moderate pain, reduce swelling, control pyrexia, and treat arthritis at higher doses [8, 9]. However, like other NSAIDs, Ibuprofen carries a risk of gastric ulcers and sometimes bleeding when taken orally.

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**MATERIALS AND METHODS**

**Materials**

Span-40, Polysorbate-60, Isooctane and methanol were purchased from Sigma-Aldrich, India. Ibuprofen was a kind gift from IOL Chemicals and Pharmaceuticals Limited India. Distilled water was used in all experiments.

**Partial phase-behavior of Sorbitan Esters/Isooctane/Water System**

Phase studies were carried out as per published literature with some modification [23] by adding double distilled water to the mixtures of isooctane and surfactant [Span-40/Polyisorbate-60 (1:1)] in different weight ratios using a magnetic stirrer. After the addition of a water drop to the mixture, the mixture was examined visually for optical clarity and through cross polarizers for the presence/absence of the birefringent liquid crystalline phase. The addition of the water was stopped when the turbidity appeared. The concentrations of surfactant, isooctane, and water were then calculated. The appearance of turbidity was chosen as an endpoint at a particular ratio of surfactant/isoctane. The phase behavior of the systems was mapped on phase diagrams, with the top apex representing the isooctane and the other apices representing the surfactant [Span-40/Polyisorbate-60 (1:1)] and water. The transparent, homogeneous, non-birefringent area enclosed within the endpoints was considered as water in oil (w/o) microemulsion. Based on the phase behavior study, OG compositions containing 1% (w/w) Ibuprofen were formulated and characterized.

**ABSTRACT**

**Objective:** This study aimed to develop and in vitro characterize an organogel (OG) loaded Ibuprofen.

**Methods:** Organogel (OG) composed of water, isooctane, sorbitan esters, sorbitan monopalmitate (Span-40), and poly(oxyethylene) sorbitan monooleate (Polysorbate-60) was loaded with Ibuprofen. The partial phase behavior of ibuprofen OG was studied to optimize the formulation composition. 1.0% w/w Ibuprofen loaded OG were characterized for rheological, in vitro release and stability study.

**Results:** Phase diagram showed an isotropic gel region at low water contents, which converted to emulsion on increasing water quantity. The rheological properties of the OG incorporating 1.0% w/w Ibuprofen shows the presence of two Tη's and elastic behavior of gel, reflects the presence of an entangled network of aqueous tubes. The fractal dimension dv value of 2.1 and 2.3 was obtained for the two curves (elastic and storage modulus), which is indicative of the formation of the densest gel structure. The diffusional release exponent (n) was found to be ~0.7 (0.5<n<1), which is indicative of non-Fickian, anomalous diffusion of the drug from the OG. The in vitro drug release exhibited release @ 7.04%/h/0.7/cm² from the OG. Ibuprofen containing OG was stable for 28 d in terms of chemical potency and gel stiffness at 4 °C and room temperature (~25 °C).

**Conclusion:** The study indicates the potential of OG for improved transdermal delivery of Ibuprofen.

**Keywords:** Microemulsion, Organogel, Phase-behavior, Gel-sol transition temperature, Fractal dimension, Ibuprofen, In vitro release, Pharmacokinetics
Gel-sol transition temperature and fractional dimension

The rheology experiments were performed as per published literature [23] using an AR500 stress-controlled rheometer (TA Instruments, U. K.). The dynamic rheology of organogel was studied by using steel parallel plate geometry of radius 20 mm and angle 0° with truncation gap 500 μm. The truncation gap was kept large to avoid breaking of the structures in the organogel sample. Frequency and strain sweep tests were carried out to specify the viscoelastic range. Sol-gel transition studies were conducted under the temperature ramp. The OG (0.16 g) was applied on the platform of the rheometer set up at a constant angular frequency (ω = 6.283 rad sec⁻¹) and the temperature was varied from 15 °C to 35 °C. The storage modulus (elastic), G', and the loss modulus (viscous), G", were recorded and graphically plotted against temperature. The sol-gel transition temperature was determined as the minima of the Tan δ (G'/G") vs temperature curve. For the determination of the fractal dimension, the rheometer was set up at a fixed temperature (20 °C) and ω was varied from 0.5 to 50 rad sec⁻¹. The relaxation exponent Δ was calculated by fitting the curve to the following power-law or a linear logarithmic equation [25].

\[ G(\omega) = \alpha_1 \omega^\alpha \text{ and } G'(\omega) = \alpha_0 \omega^\alpha \] .............................. [1]

\[ \log G(\omega) = \log \alpha_1 + \alpha \log \omega \text{ and } \log G'(\omega) = \log \alpha_0 + \alpha \log \omega \] .............................. [2]

where G(ω) is the modulus, ω is the angular frequency (rad sec⁻¹), α is a constant and Δ is the relaxation exponent. After the determination of relaxation exponent, a relationship of relaxation exponent with the fractal morphology of the gel network was established as published by Muthkumar [26] in the following equation:

\[ \Delta = (d + 2 - 2d_f)/2(d + 2 - d_f) \] .............................. (3)

where d is dimension and d_f is the fractal dimension. For gel, d = 3, therefore the equation could be simplified as:

\[ d_f = (15 - 10d)/6 - 2d \] .............................. (4)

Drug release through synthetic membrane

The Ibuprofen containing OG was subjected to an in vitro drug release study through a cellophane membrane. The formulation (1.0g) was taken in a pretreated dialysis bag (as per the procedure suggested on the supplier, Sigma-Aldrich) clipped on both sides. The dialysis bag was immersed in 100 ml phosphate-buffered saline (PBS, pH 5.5) in the beaker maintained at 37±1 °C on a temperature-controlled, magnetic stirrer plate. The contents of the beaker were stirred at 50 rpm using the magnetic bar. The sample from the beaker was withdrawn periodically for 24h and replaced immediately with an equal volume of PBS, pH 5.5. The samples were analyzed as explained later. The mass released (M) with time (t) was calculated according to the following power law [27].

\[ \frac{M}{M_\infty} = t^n \] .............................. [5]

where \(M_\infty\) is mass released after an infinite time.

Stability

The OGs containing the drug and respective placebos were stored for 15, 30, and 45d at 4 °C and 25 ± 5 °C in collapsible aluminum tubes. The drug content was determined spectrophotometrically after dissolving the formulation in methanol against similarly treated placebos. Viscosities of OG formulations were also determined at weekly intervals using a Brookfield Digital Viscometer model LVDV-I+ (Brookfield Engineering Labs. Inc., USA). The formulations were also subjected to mechanical stress by centrifuging (Remi Centrifuge, India) at 10000 rpm for 20 min and the volume of phase separation of OGs was noted.

RESULTS AND DISCUSSION

Phase behavior of OGs

Phase behavior of OGs system consisting of isooctane, water, and surfactant mixture (Span-40 and Polysorbate-60) was investigated by plotting the pseudo-ternary phase diagrams in which each corner of the diagram represents 100% of that particular component. Fig. 1 shows the phase diagram of the isooctane, water, and surfactant mixture. From the phase diagrams, it is clear that the area of OGs region decreased with the addition of the drug (grey). This correlates well with a previous study [23].

\[ G(\omega) = \alpha_1 \omega^\alpha \text{ and } G'(\omega) = \alpha_0 \omega^\alpha \] .............................. [1]

\[ \log G(\omega) = \log \alpha_1 + \alpha \log \omega \text{ and } \log G'(\omega) = \log \alpha_0 + \alpha \log \omega \] .............................. [2]

Fig. 1: Triangular phase diagram of OG formulations containing surfactant (Span40/Polysorbate-60, 1:1), isooctane and water.

The encapsulated gray area in dark area (without Ibuprofen) shows the phase behavior of ibuprofen containing OG

Rheological characteristics of OGs

Gel-sol transition temperature (Tg) of OG formulations was determined using a Rheometer set up at a constant angular frequency ω = 6.283 rad sec⁻¹ and the temperature, T, were varied from 15 °C to 35 °C. A representative loss of storage modulus (G') and loss modulus (G") versus temperature curve obtained during the gelation process was plotted as shown in fig. 2. The curves present three domains: high and nearly constant values of G' corresponding to sol state, a sharp increase corresponding to gel microstructure formation and the last domain with a tendency to a plateau that could be linked to gel formation. Both the G' and G" values were slightly increased with an increase in drug concentration at a particular temperature. The Tan δ was calculated as G"/G'. The curve between Tan δ and T was plotted and the Tg is obtained from the minima of the curve (fig. 2, inset). The parameter δ is the phase angle or phase shift between the deformation and response that is measured. It is the point where the viscoelasticity of the gel changes abruptly. A careful observation into this curve shows the presence of a shoulder (indicated by an arrow), indicating that the presence of two Tg's. Thus, gel-sol conversion can be explained by a change in the arrangement of the gel network in two stages due to the incorporation of the drug, which suggests that the type of drug encapsulated in organogel affects OG network.

The viscoelastic nature of gels is a function of the microstructure network, which can be evaluated using the rheometer set up at fixed temperature (20 °C) and varying ω from 0.5 to 100 rad sec⁻¹. The elastic and viscous modulus shows an asymptotic increase in values, with the elastic modulus component [G'(ω)] always having a higher value than the viscous modulus component [G"(ω)], suggesting the predominately elastic nature of the sorbitan ester OG within the indicated frequency range. The elastic behavior, in this case, reflects the presence of an entangled network of aqueous tubules. Fractal morphology of OG could be determined from the fractal dimension (d_f) through a theoretical model for polyelectrolyte gels [25]. The relaxation exponent, Δ, was calculated fitting the power law equation to the G'(ω), G"(ω) vs curves or by plotting logarithm [G'/(ω)], G"/(ω)] vs logarithm [ω] curves and determining the slope from the linear fit equation (fig 3). The d_f value of 2.1 and 2.3 was obtained for the two curves (elastic and storage modulus), which is indicative of the formation of the densest gel structure. The values corroborate with reported studies [23].
Drug release from OGs

The kinetics of ibuprofen release from OG was established by plotting a graph between percent cumulative drug release against time and the value of the diffusional release exponent (n) was calculated by fitting a power law to the curve (fig. 4). The value of n was found to be ~0.7 (0.5<n<1), which is indicative of non-Fickian, anomalous diffusion of the drug from the OG [27]. A graph of the percent cumulative amount of drug vs time 0.7 gives the release rate from its slope. The *in vitro* drug release exhibited release @ 7.04%/h 0.7/cm² from the OG.

Stability

The results of stability studies on ibuprofen containing OG formulation are summarized in table 1. The assay of drugs was more than 98% in all the formulations at the end of 45 d at 4 °C and room temperature (~25 °C) indicating that the chemical stability of ibuprofen in OG formulations was good. The formulations became stiffer over a 45 d period which was shown by an increase in the viscosity of the OG formulations. The OG formulations were resistant to centrifugal stress.

### Table 1: Stability of ibuprofen OG formulation

| S. No. | Formulation code | Time  | Assay (%) | Viscosity (cps) | Centrifugal Stress* |
|--------|------------------|-------|-----------|----------------|---------------------|
| 1.     | Assay (%)        | 15d   | 99.1±1.2  | 1200           | nil                 |
|        |                  | 30d   | 98.4±1.4  | 1600           | nil                 |
|        |                  | 45d   | 98.7±0.9  | 2000           | nil                 |

n=3 (Mean of three determination) *Phase separation on the centrifugation at 1000 rpm for 20 min

CONCLUSION

Organogel was made using of sorbitan monopalmitate/polysorbate, isooctane and water. The sol-gel transition with decreasing temperature shows an abrupt denser network formation indicated by the higher fractal dimension value. The drug permeation was slow and displayed anomalous release behavior. The storage stability of OG was good for 45 d. This study proves that sorbitan monopalmitate/polysorbate-based organogel can be a good drug delivery system for transdermal delivery.

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AUTHORS CONTRIBUTIONS

All the authors have contributed equally.

CONFLICT OF INTERESTS

Declared none

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