ANTI-EPILEPTIC DRUG LOADED NIOSOMAL TRANSDERMAL PATCH FOR ENHANCED SKIN PERMEATION

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ABSTRACT

Objective: To formulate and characterize midazolam loaded niososomal transdermal patches for overcoming the frequent dosing and lower bioavailability complications associated with conventional therapy.

Methods: The loaded niososomal transdermal patches were prepared by thin film hydration method. The preliminary evaluation and characterization studies were conducted to find the optimized formulation. The in vitro release and ex vivo permeation studies were investigated. The histopathological studies and stability studies were also assessed.

Results: The midazolam loaded niosomal transdermal patches of vesicle size and zeta potential 116.1±84.46 d. nm and 8.56±1.2 mV respectively was formulated. The characterizations of both niosome and niosomal transdermal patches were found to be within the acceptable limits. The in vitro drug release showed an initial burst release followed by sustained release for both optimised niosomal formulation N5 and optimised niosomal transdermal patch formulation NT5 with a maximum activity at 97.3±0.35% and 98.9±0.20% respectively. The ex vivo permeation studies of niosomal transdermal patch NT5 was performed which showed a higher permeability than control solution with a flux value of 0.151. The histopathological studies of the optimised formulation showed no detectable lesions upon skin surface and irritations. The stability studies showed that patches were stable over 90 d in different atmospheric conditions.

Conclusion: The midazolam loaded niosomal transdermal patch was found to be a promising nano drug delivery alternative which showed better entrapment, release with permeation profile for the daily management of epilepsy with decreased dosing frequency.

Keywords: Status epilepticus, Midazolam, Niosome, Soya bean oil, Niosomal transdermal patch, Gelatin

INTRODUCTION

Epilepsy is a disorder of the brain characterised by an enduring predisposition to generate epileptic seizures and by neurobiologic, cognitive, psychological and social consequences of this condition [1]. The epilepsy associated seizures are distinct to a person and undetectable sometimes and thus require special attention. Seizure is an event of transient occurrence of signs or symptoms due to abnormal, excessive or synchronous neuronal activity of the brain which can vary from short to long periods of vigorous shaking [2]. The excited electrical activity underlying epilepsy is due to biochemical processes at the cellular level promoting neuronal hyperexcitability. Any abnormalities in the normal neuronal transmissions, starting from a single enzyme-receptor abnormality to several key cortical and subcortical structures are involved in generating clinical seizures [3]. Seizures arise from an excessively synchronous excitation and sustained discharge of a group of cortical neurons. Prevailing seizures tend to recrudescence and have no immediate underlying genesis which are the result of excessive and abnormal nerve cell activity in the cortex of the brain [4–6]. Many currently used medications, even intravenous preparations are found to be least effective in early stages of epilepsy treatment [7]. The currently used major formulations for treatment of epilepsy include oral dosages which were once promising approaches for treatment methods but were less efficient in its therapeutic effect [8]. Novel drug delivery research provide a wider opportunity for these limitations [9–11]. Niosomes are nanometre size hydrated vesicles composed of non-ionic lamellar lipid bilayer posing an amphiphilic infrastructure [12–15]. The drug gets encapsulated within the amphiphilic system of the biodegradable and biologically compatible carrier system [16]. The novel carrier molecules are formulated by admixture of non-ionic surfactants like Span 60 or from dialkylpolyglycerol ether class which is stabilised by incorporating waxy steroids like cholesterol [17–20]. The niosomes are already leading in the market where they are used to incorporate various anti-cancer, anti-inflammatory drugs as well as hormonal preparations [21]. Niosomes possess advantages than conventional liposomes and tackles the susceptibility of phospholipids to undergo oxidative degeneration and variations in quality of lipids [22]. The transdermal route of drug administration ensures systemic delivery of drug by applying a drug formulation onto intact and healthy skin thus ensuring sustained drug release and bypass of first-pass metabolism [23–26]. Transdermal drugs significantly delivers molecules in a potent quantity that overcome the conventional problems of oral dosing. Thus the aim of the study was to develop midazolam loaded niosomal transdermal patches an effective substitute for the existing maintenance therapies used for controlling epileptic seizures [27].

MATERIALS AND METHODS

Materials and excipients

The drug of choice, midazolam, was provided as a gift sample by Lake Chemicals Private Limited, Bangalore. Soya bean oil was obtained, fromSigma Aldrich. All other chemicals used in the experiment were of analytical grade.

Preformulation studies

Fourier transform infrared (FTIR) spectroscopy

The FTIR spectra of drug and excipients were obtained to ascertain the compatibility between midazolam and selected polymers using FTIR spectrophotometer by KBf pellet method [28].

Solubility studies

Solubility studies of drug were determined in different solvents such as distilled water, ethanol, methanol, isopropyl alcohol, acetone, n-butyl alcohol and phosphate buffer pH (5.5).

Melting point of the drug

The melting point of the sample was determined by open capillary method [29].

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Partition coefficient

The partition coefficient of midazolam in n-octanol was found out and the concentration of drug content in the aqueous layer is determined spectrophotometrically [30].

Absorption maxima of midazolam in methanol and phosphate buffer solution (PBS)

The absorption maxima of midazolam in both 10 ml of methanol and PBS pH 5.5 was found to be 220 nm using UV-Vis spectrophotometer [31].

Drug compatibility with excipients

The compatibility of the drug with excipients is determined using FTIR spectral analysis [32].

X-Ray diffraction studies

XRD pattern was documented for midazolam API, optimised niosome NS and optimised niosomal patch NT5 on Joel JDX 8030 X-ray diffractometer (MTI Corporation, USA) for a specified quantity of each preparation [33, 34].

Formulation of midazolam niosome and transdermal patch preparation

The niosomal dispersion of midazolam was prepared by thin film hydration method using rotary vacuum evaporator. Surfactant (span60), soya bean oil, cholesterol in ratio 2:1:0.75 was weighed and dissolved together in the round bottom flask containing methanol-chloroform mixture of ratio 2:1 together with glass beads. The round bottom flask is connected to a rotary vacuum evaporator and then vacuum was applied. This round bottom flask was rotated at 150rpm and bath temperature was set at 28 °C. In this first phase, the vacuum was applied to completely remove the solvents until the complete dried solid thin film was developed in the wall of the flask. The film was hydrated by adding PBS solution pH 5.5 containing the dissolved drug. The hydrated solution was again agitated for 1 hr at 60 °C for complete hydration. The obtained niosome dispersion was sonicated using probe sonicator for 20 min to decrease the particle size. The obtained mixture was stirred for 1 hr in a magnetic stirrer to obtain a uniform niosome [35-37].

The obtained niosomes were stored under normal room temperature and transdermal patches were prepared using solvent casting method. The polymer solution of gelatin was prepared in a 250 ml beaker by adding 20 ml of distilled water at 35 °C. The 1% PEG 4000 and 0.1% turpentine oil was added as permeation enhancers. To this the prepared niosomal dispersion was introduced (span60), soya bean oil, cholesterol in ratio 2:1:0.75 was weighed and dissolved together in the round bottom flask containing methanol-chloroform mixture of ratio 2:1 together with glass beads. The round bottom flask is connected to a rotary vacuum evaporator and then vacuum was applied. This round bottom flask was rotated at 150rpm and bath temperature was set at 28 °C. In this first phase, the vacuum was applied to completely remove the solvents until the complete dried solid thin film was developed in the wall of the flask. The film was hydrated by adding PBS solution pH 5.5 containing the dissolved drug. The hydrated solution was again agitated for 1 hr at 60 °C for complete hydration. The obtained niosome dispersion was sonicated using probe sonicator for 20 min to decrease the particle size. The obtained mixture was stirred for 1 hr in a magnetic stirrer to obtain a uniform niosome [35-37].

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Scanning electron microscopy

The scanning electron microscope (SEM) was used to scan the atomic surface of midazolam niosomes and niosomal transdermal patches which was view under electron microscope for shape and size [43, 44].

In vitro release studies

The in vitro release studies of different niosomal formulations of midazolam was done by static dissolution method. The apparatus was set up by introducing an open end tube aligned vertically to a beaker containing the 50 ml of PBS pH 5.5 which acts as receptor compartment. The open end tube was tied with a cellophane membrane on one side which resembles the membrane acting as a barrier within the body, so that only the tip touches the surface of buffer solution. The reaction conditions were standardised throughout the study. The receptor compartment was filled with PBS pH 5.5 and the donor compartment with 1 ml vesicular dispersion of niosomal midazolam. The beaker was placed on a magnetic stirrer for a definite study time and samples were withdrawn at regular intervals. For each withdrawal the fresh buffer was replaced into the beaker and collected samples were diluted using pH 5.5. The percentage of drug release in the receptor medium was spectrophotometrically analysed using UV-visible spectrophotometer at 220 nm [45].

Kinetic model of in vitro drug release studies

The in vitro release studies data were fitted to various kinetic models like Zero order, First order, Higuchi Plot and Koresmeyer peppas plot to identify the model of drug release [46, 47].

Ex vivo permeation study

The permeation studies was done in Franz diffusion cell where the mucosal tissue was kept at the donor compartment and 1 ml of the formulation in the receptor compartment. With a needle, required aliquots at regular intervals were transferred to sample bottles and was replaced with 1 ml fresh buffer. The samples were suitably diluted and spectrophotometrically analysed by UV-Vis spectrophotometer at 220 nm. The procedure was done for continuous 8 h and the optimised niosome formulation, optimised niosomal patch and drug in buffer PBS pH 5.5 was used for the study [48-50].

Characterisation of niosomal transdermal patch

Flatness

The flatness test was conducted circular patches prepared which were cut into three different longitudinal strips and measured for uniformity in flatness [51].

Swelling ratio

The swelling properties of transdermal patches were found by placing completely dried patches in a beaker containing 250 ml of PBS pH 5.5 and maintained it at 27 °C. After 24 h the patches were taken and measured the water uptake using a digital balance [52].

Weight uniformity

The uniformity in weight of different transdermal patch formulations was done by separately weighing the selected patches individually calculated for its average.

Thickness

The thickness of transdermal patches was measured using screw gauge at different sites of the various formulations and average measurement of triplicate readings was taken [53].

Percentage drug content

1 cm sized patches were cut out and dissolved in vortex shaker containing 5 ml of methanol for 5 min. The solution was made up to the volume by 10 ml methanol and spectrophotometrically analysed for percentage drug content.

Percentage moisture content

The transdermal patch was introduced to a desicator at room temperature containing activated silica for one day. The patches
were repeatedly weighed the next day for constant weight was attained [54].

**Folding endurance**

A longitudinal strip of a transdermal patch was cut and folded repeatedly and calculated the number of times the patch was folded at the same place without breaking/rupturing [55].

**Transmission electron microscopy (TEM)**

The optimized midazolam niosomal transdermal patch NT5 was viewed under TEM where the electron beams were used to emit radiation through the sample to form an image [56-58].

**Histopathological studies**

**Source of porcine ear skin and its preparation**

The ear of the freshly sacrificed pig was collected from a local slaughter house at morning and was cleaned with distilled water to be used for the study by noon, housed at 23-27 °C. The underlying fat deposits were initially excised off uniformly and the skin layers from the outer. The skin preparation retaining all the layers was allowed to equilibrate for one hour in receptor buffer to retain its properties as live tissue. Later on the skin was stored in containers partly filled with formalin solution.

**Preparation of formalin solution**

Firstly 100 ml of formalin was solubilised in distilled water and to it disodium hydrogen phosphate and sodium hydrogen phosphate was added and pH was made up to 6.6.

**Preparation of porcine ear skin for histological studies**

The prepared pig skin was mounted between the compartments of Franz diffusion cell for ex vivo permeation studies. The histopathological studies of the treated skin were done for finding observable changes in tissue characteristics. For the study the optimised niosomal formulation, N5 and niosomal transdermal patch, NT5 was treated on control ear tissue and after fixation was viewed under the light microscope under standard atmospheric conditions [59-61].

**Stability studies**

The optimised niosomal formulations of niosome and transdermal patch were subjected to various stability studies for 45 d within refrigerator (4±2 °C) and room temperature (29±2 °C) in triplicate [62].

**RESULTS AND DISCUSSION**

**Preformulation studies**

**Fourier transform infrared (FTIR) spectroscopy**

The FTIR of an obtained pure drug of midazolam and midazolam loaded niosomal transdermal patch was found to be in accordance with monograph (IP) as shown in fig. 1(A) and fig. 2(B) [63].

**Solubility studies**

The pure drug is partially insoluble in distilled water and n-butyl alcohol and completely soluble in methanol, acetone, ethanol, isopropl alcohol, PBS pH 5.5.

**Melting point of the drug**

The melting point of the drug was found to be 158-160 °C and it was in accordance with that of the reference.

**Partition coefficient**

The partition coefficient of drug midazolam was found to be 4.5 and shows an amphiphilic character when entering body pH.

**Formulation of midazolam loaded niosomal transdermal patch**

The niosome of midazolam was formulated using thin film hydration method. The surfactants, Span 60 and soya bean oil (natural surfactant) with co-surfactant, cholesterol was used for formulating niosome using rotary vacuum evaporator. 2%w/v of span 60, 1%w/v of cholesterol and 0.75%w/v soya bean oil had the ability to convert within itself to a thin film in methanol chloroform solvent system. 2%w/v of span 60 acts as a surfactant and stabilises the chemical attachments between the bilayer of niosome with cholesterol. The 1%w/v of cholesterol acts as co-surfactant and stabilises the biomembrane along with span 60 [64]. The 0.75%w/v soya bean acts as a natural surfactant and emulsifies or stabilises the niosomal membranes [65-66]. Methanol and chloroform were used as solvents to reconstitute the contents of the thin film obtained from the round bottom flask of rotary vacuum evaporator.

The transdermal patch was prepared using solvent casting method using gelatin as natural polymer [67]. Gelatin 2%w/v was used to get the optimised transdermal patch formulation were gelatin is used for its excellent natural film-forming properties [68-69]. The prepared niosomes were loaded into the patch formulation to which turpentine oil was added as a natural permeation enhancer [70]. The formulations of niosomal transdermal patches are shown in table 1(A) and table 1(B).

### Table 1(A): Formulation composition of midazolam loaded niosome

<table>
<thead>
<tr>
<th>S. No.</th>
<th>Formulation code</th>
<th>Midazolam (mg)</th>
<th>Span 60 (mg)</th>
<th>Cholesterol (mg)</th>
<th>Soya bean oil (%)</th>
<th>Methanol (%)</th>
<th>Chloroform (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1)</td>
<td>N1</td>
<td>1</td>
<td>1</td>
<td>0.5</td>
<td>0.75</td>
<td>3</td>
<td>2</td>
</tr>
<tr>
<td>2)</td>
<td>N2</td>
<td>10</td>
<td>1</td>
<td>1</td>
<td>0.25</td>
<td>3</td>
<td>2</td>
</tr>
<tr>
<td>3)</td>
<td>N3</td>
<td>10</td>
<td>1</td>
<td>1.25</td>
<td>0.5</td>
<td>6</td>
<td>4</td>
</tr>
<tr>
<td>4)</td>
<td>N4</td>
<td>10</td>
<td>1.75</td>
<td>1.25</td>
<td>1</td>
<td>8</td>
<td>6</td>
</tr>
<tr>
<td>5)</td>
<td>N5</td>
<td>10</td>
<td>2</td>
<td>1</td>
<td>0.75</td>
<td>10</td>
<td>8</td>
</tr>
<tr>
<td>6)</td>
<td>N6</td>
<td>10</td>
<td>2.5</td>
<td>1.5</td>
<td>1</td>
<td>9</td>
<td>5</td>
</tr>
<tr>
<td>7)</td>
<td>N7</td>
<td>10</td>
<td>3</td>
<td>1</td>
<td>0.75</td>
<td>8</td>
<td>6</td>
</tr>
</tbody>
</table>

### Table 1(B): Formulation composition of midazolam loaded niosomal transdermal patches

<table>
<thead>
<tr>
<th>S. No.</th>
<th>Formulation code</th>
<th>Midazolam loaded niosomes N5 optimised formulation (ml)</th>
<th>Gelatin (mg)</th>
<th>Distilled water qs.20</th>
<th>PEG 4000 (%)</th>
<th>Turpentine oil (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1)</td>
<td>NT1</td>
<td>1</td>
<td>6</td>
<td>q.s.20</td>
<td>5</td>
<td>4</td>
</tr>
<tr>
<td>2)</td>
<td>NT2</td>
<td>1</td>
<td>8</td>
<td>q.s.20</td>
<td>2</td>
<td>2</td>
</tr>
<tr>
<td>3)</td>
<td>NT3</td>
<td>1</td>
<td>5</td>
<td>q.s.20</td>
<td>0.1</td>
<td>1</td>
</tr>
<tr>
<td>4)</td>
<td>NT4</td>
<td>1</td>
<td>4</td>
<td>q.s.20</td>
<td>3</td>
<td>0.05</td>
</tr>
<tr>
<td>5)</td>
<td>NT5</td>
<td>1</td>
<td>2</td>
<td>q.s.20</td>
<td>1</td>
<td>0.1</td>
</tr>
<tr>
<td>6)</td>
<td>NT6</td>
<td>1</td>
<td>1</td>
<td>q.s.20</td>
<td>2</td>
<td>0.25</td>
</tr>
<tr>
<td>7)</td>
<td>NT7</td>
<td>1</td>
<td>3</td>
<td>q.s.20</td>
<td>4</td>
<td>5</td>
</tr>
</tbody>
</table>
Lamda max of midazolam in methanol and phosphate buffer solution (PBS)
The absorption maxima of midazolam in both 10 ml of methanol and PBS pH5.5 was found to be 220 nm using UV-Vis spectrophotometer.

Drug compatibility with excipients
The FTIR spectrum of all individual excipients was compared with the FTIR spectrum of the two formulations. The drug, as well as excipients, exhibited no changes in its characteristic peaks when loaded to niosomal and transdermal patch formulation.

X-ray diffraction studies
The XRD pattern of drug midazolam, N5 niosomal formulation, and NT5 niosomal transdermal patch formulation are show in fig. 1(C) to fig. 1(E). The characteristic intense and sharp peaks in the XRD pattern of pure midazolam depicts the crystalline nature, appearing at 9.5, 11.7, 13.3, 20.7, 22.7 2-Theta values. During the formulation of niosomes, the number of intense peaks lowered and confined to 5.3 and 21.6 2-Theta values. The XRD pattern of niosomal transdermal patch formulation showed only one intense peak at 5.1 2-Theta values. While comparing the results of drug with optimized formulations, the concluding pattern shows a lower intensity and no. of peaks. This apparently illustrates that the extent of crystalline nature of midazolam was reduced when the drug was formulated as niosome and further loaded to a transdermal patch formulation.

Characterisation
Characterisation of niosomes
Vesicle size
The particle size distribution of midazolam niosomes assessed using DLS showed an average particle size of 116.1±84.46 d. nm. The size was analysed to be sufficiently small to penetrate the pores of stratum corneum as shown in fig. 2(A).

![Fig. 1: (A) FTIR Spectrum of midazolam loaded niosome, (B) FTIR Spectrum of midazolam loaded niosomal transdermal patch, (C) X-ray diffraction studies of pure drug, (D) X-ray diffraction studies of optimised N5 formulation,(E) X-ray diffraction studies of optimised NT5 formulation](image-url)
Zeta potential

In order to analyse the approach of all niosomal midazolam formulations and their ionic interaction with the biological membrane, the zeta potential was analysed to be 8.56±1.2 mV.

The zeta potential directly indicates the surface charge of the lipid nanocarrier. Due to its positive value, the lipid carrier interacts with the negatively charged biological membrane so as to increase the cellular uptake [71]. Thus the zeta potential of a stable dispersion of niosomes is shown in fig. 2(B).

Entrapment efficiency

The entrapment efficiency of all niosomes ranged between 88.79±0.21% to 90.15±0.34% as shown in fig. 2(C). The maximum entrapment was found to be 90.15±0.34% for the niosome formulation N5. The entrapment efficiency of niosomes was dependent on the concentrations of polymers used [72]. By increasing the concentration of polymers, multiple coating layers will be formed around the nanoparticle which will retard drug release [73].

SEM

The SEM studies were done to examine the morphological shape of the niosomal midazolam. The SEM images for the midazolam niosome and niosome loaded transdermal patches are shown in fig. 2(D) and fig 2(E) which confirms the smooth spherical nature of niosomes.

TEM

The optimised niosomal transdermal patch NT5 was examined to TEM studies were the niosomal particles were found to be nearly spherical. The size of the particles increased according to the molecular weight of gelatin [74]. The image depicted clear intact particles with the least aggregate formation even after loading niosome into the transdermal patch as seen in fig. 2(F).

In vitro release studies

The in vitro release studies of various formulations of midazolam loaded niosome was done for 8 h in acetate buffer pH 5.5. A constant temperature and pressure conditions were maintained throughout the experiment. The drug release was found to be maximum by N5 formulation with the highest release of 97.3±0.35%. The excipients used in the preparation of niosomes in a proportional concentration
played a major role in the release characteristics [75-76]. The in vitro drug release studies of midazolam niosomal formulations at different time intervals were shown in fig. 3.

**Kinetic model of in vitro drug release studies**

The drug release profile of N5 niosomal midazolam with highest drug release pattern was attributed to different kinetic models like Zero order model, First order Model, Higuchi's diffusion model, and Koresmeyer peppas plot to interpret drug release by kinetic modelling [77]. The release kinetics of drug was found to be Zero order as the $R^2$ regression coefficient of the model was found to be 0.9597. This was best fitted with Koresmeyer peppas plot ($n = 0.4201$) which is greater than 0.5 depicting non fickian transportation mechanism as depicted in table 2 [78].

![In vitro Drug Release Studies](image)

**Fig. 3: In vitro release studies of different midazolam niosomal formulations (values are expressed as mean±standard deviation, n=3)**

<table>
<thead>
<tr>
<th>Table 2: Kinetic models of optimised N5 niosomal formulation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Zero-order release plot</td>
</tr>
<tr>
<td>$R^2$</td>
</tr>
<tr>
<td>0.9597</td>
</tr>
</tbody>
</table>

**Characterisation of niosomal transdermal patch**

**Flatness**

The different formulations of midazolam loaded niosomal transdermal patch was measured in triplicate and there were slight variations in flatness between the different formulations indicating no significant stretching or constriction of the patches. The NT5 formulation showed highest uniformity in flatness as shown in fig. 4(A).

**Swelling ratio**

The swelling studies of all formulations were found to be a function of polymer concentration. The % swelling ranged from 22.96±0.03 to 38.95±0.03 as seen in fig. 4(B). It was seen that the polymer concentration varied with the amount of water it could absorb during the study and was found ideal for N5 formulation [79]. The swelling of the polymer was due to the dissolution of gelatin in the buffer as a function of temperature and time.

**Weight uniformity**

The average weight of different transdermal patch formulations ranged from 152.35±0.02 to 168.46±0.01 mg as seen in fig. 4(C). All the formulations were found to be having satisfactory results and has low standard deviation values within a formulation.

**Thickness**

The thickness of various transdermal patches prepared varied between 0.262±0.012 to 0.315±0.023 mm as shown in fig. 3(D). The difference in thickness showed very less standard deviation hence the formulations did not show any susceptible changes [80].

**Percentage of drug content**

The drug content of different transdermal patches was found to be 90.26±0.071 to 90.52±0.073 as shown in fig. 4(E). The values show least deviations depicting the uniform distribution of drug content throughout the patch. The drug retained within the patch was comparatively high for N5 formulation.

**Percentage moisture content**

The prepared patches showed minimum percentage moisture content ranging between 0.002±0.002 % to 0.0038±0.004 % as shown in fig. 4(F). The lower concentration of moisture uptake helps in maintaining the preparation free from microorganism's contamination [81]. The results ensures the maintenance of proper architecture and stability of patches which was found to be least within N5 formulations.

**Folding endurance**

The folding endurance of the patches was reported to be 234±6.36 to 271±3.54 as shown in fig. 4(G). The patches N1 and N5 formulations having an intermediate concentration of gelatin was found to be having the satisfactory results. This ability to retain the structural integrity helps the patch to be retained over the skin surface for a longer time without breaking [82, 83].

**In vitro release studies**

The in vitro release studies of various formulations of midazolam loaded niosomal transdermal patch was also done for 8 h in phosphate buffer pH5.5. A constant temperature and pressure conditions were maintained throughout the experiment. The drug release was found to be maximum by NT5 formulation with the highest release of 98.9±0.20%thus regarded as the optimised formulation. The release was initially found to be burst followed by the gradual sustained increase in free drug concentration. The hydrophilic polymer gelatin absorbs fluid and enables the significant swelling of the patch [84]. Thus along with plasticizers like PEG 4000, it enhances the drug release with improved and flexible characteristics [85-86]. The in vitro drug release studies of midazolam niosomal transdermal patch at different time intervals are shown in fig. 5.
Kinetic model of in vitro drug release studies

The drug release profile of midazolam loaded niosomal transdermal patch (NT5) with highest drug release pattern was attributed to different kinetic models like Zero order model, First order Model, Higuchi's diffusion model, and Koresmeyer peppas plot to interpret drug release by kinetic modelling. The release kinetics of drug was found to be Zero order as the \( R^2 \) regression coefficient of the model was found to be 0.9565. This was best fitted with Koresmeyer peppas plot \( (n = 0.4953) \) which is greater than 0.45 depicting non fickian transportation mechanism. The kinetics also obeys the Higuchi plot were \( R^2 \) value is almost 0.9585 as illustrated in table 3.

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**Fig. 4:** (A) Flatness, (B) Swelling ratio, (C) Weight uniformity, (D) Thickness, (E) Percentage drug content, (F) Percentage moisture content, (G) Folding endurance of different niosomal transdermal patch

**Fig. 5:** In vitro release studies of different midazolam loaded niosomal transdermal patch formulations, (values are expressed as mean±standard deviation, \( n=3 \))
Table 3: Kinetic models of optimised NT5 niosomal transdermal patch formulation

<table>
<thead>
<tr>
<th></th>
<th>Zero order release plot</th>
<th>First order release plot</th>
<th>Higuchi plot</th>
<th>Koresmeyer peppas plot</th>
</tr>
</thead>
<tbody>
<tr>
<td>$R^2$</td>
<td>0.9565</td>
<td>0.9019</td>
<td>0.9585</td>
<td>0.4953 0.9639</td>
</tr>
</tbody>
</table>

**Ex-vivo permeation studies and steady state flux value determination**

The permeation studies were done in Franz diffusion cell with normal pig skin mucosa for 8 h. The mucosa biologically resembles human skin mucosa and thus ex vivo studies were done with pig skin [87]. The pigskin mucosa was experimented with the optimized niosomal formulation N5, optimised niosomal transdermal patch NT5 and buffer containing drug preparation (Control). The samples withdrawn at each intervals were analysed and the cumulative amount of drug permeated v/s time was plotted. The permeation was found to be enhanced in case of niosomal midazolamN5 and when the niosome was loaded to transdermal patch NT5 in a controlled manner from the porcine skin. The permeation is sustainably enhanced for formulations with an optimum concentration of Span 60 due to its increased wettability, drug distribution and ability to lower the surface tension [88]. The natural oil, soya lecithin provides an improved penetration effect by increasing its permeability into the carrier vesicles and thereby decreasing the resistance of molecules by enhancing the fluidity [89-90]. The flux values of optimised formulations reported the drug penetrated during each hour per square centimeter and established a relation between an amount of drug released and swelling of patch [91]. The ex vivo permeation studies of the above mentioned 3 formulations is given in fig. 6. The values of permeation parameters (steady state flux (J), permeation coefficient (P) and enhancement ratio of optimized midazolam loaded niosomal transdermal patch (NT5), optimised midazolam loaded niosome (N5), midazolam API loaded transdermal patch and drug in PBS pH 5.5 (control drug solution) was illustrated in table 4.

![Ex vivo permeation comparison of selected formulations with control drug solution](image)

Table 4: Flux, enhancement, and permeation coefficient of optimized niosomal midazolam transdermal patch NT5

<table>
<thead>
<tr>
<th>S. No.</th>
<th>Parameters</th>
<th>Midazolam loaded niosomal transdermal patch (NT5)</th>
<th>Midazolam loaded niosome (N5) (*from previous study)</th>
<th>Midazolam API loaded transdermal patch</th>
<th>Drug in PBS pH 5.5 (control drug solution)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.</td>
<td>Steady state flux-J (µg/cm²/h)</td>
<td>4.26±0.31</td>
<td>1.13±0.22</td>
<td>0.62±0.24</td>
<td>0.56±0.14</td>
</tr>
<tr>
<td>2.</td>
<td>Enhancement ratio (basis of control drug solution)</td>
<td>7.61</td>
<td>2.01</td>
<td>1.11</td>
<td>-</td>
</tr>
<tr>
<td>3.</td>
<td>Enhancement ratio (basis of midazolam API loaded transdermal patch)</td>
<td>6.84</td>
<td>1.81</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>4.</td>
<td>Permeation coefficient- Kp(cm²/h)</td>
<td>0.0062±0.47</td>
<td>0.0031±0.33</td>
<td>0.007±0.28</td>
<td>0.56±0.42</td>
</tr>
</tbody>
</table>

(Values are expressed as mean±standard deviation, n=3)

The flux value obtained for optimised midazolam loaded niosomal transdermal patch (NT5) was found to be 4.26 µg/cm²/h and permeation coefficient of 0.0062 cm²/h. These values correlate that optimised midazolam loaded niosomal transdermal patch (NT5) showed greater steady state flux values when compared to midazolam loaded niosome (N5) midazolam API loaded transdermal patch and control drug solution. The ex vivo permeation parameters confirms that the permeation of midazolam loaded niosomes across the porcine skin barrier is significantly higher when compared to the drug in PBS pH 5.5 (control drug solution). The primary reason for niosomes to enhance the permeation of drugs is through the structural modification in stratum corneum layers. Due to the effect of natural permeation enhancers and non-ionic surfactants used in the formulation of niosomes, the intercellular lipid barrier gets dramatically loosened. The enhanced permeability may also occur due to aggregation and fusion of midazolam loaded niosome at the interface of stratum corneum and generation of high thermodynamic activity of midazolam which was highly concentrated in the bilayers. As a result, enhanced flux is achieved due to the direct transfer of drug from vesicles to the skin. The fig. 7 shows the comparison of steady state flux of selected formulations through porcine skin layers after ex vivo permeation studies.
Fig. 7: Comparison of steady-state flux of selected formulations through porcine skin layers (Values are expressed as mean±standard deviation, n=3)

**Statistical analysis by student t-test**

The statistical analysis by student t-test was performed for the steady state flux values of different formulations. The statistical analysis by student t-test revealed that there is significant difference in the steady state flux value of the optimised midazolam loaded niosomal transdermal patch (NT5) and optimised midazolam loaded niosome (N5) with that of drug in PBS pH 5.5 (control drug solution), where (P<0.05) thus very significant. Hence the difference is statistically significant.

Fig. 8: Histopathological evaluation of porcine skin. 8(A) Porcine skin treated with midazolam loaded niosome N5, 8(B) Porcine skin treated with midazolam loaded niosomal transdermal patch, 8(C) porcine skin treated with control (Drug in buffer solution), 8(D) Normal porcine skin
Histopathological examination
The porcine mucosa subjected to ex vivo permeation studies were assessed for histopathological changes [92]. The fig. 8(A),(B),(C) shows that neither the optimized midazolam loaded niosome (N5), optimised midazolam loaded niosomal transdermal patch (NT5) and drug loaded in buffer solution PBS pH 5.5, shows no significant changes in the histological pattern when compared to normal mucosadepicted in fig. 8 (D). Hence the midazolam loaded niosomal transdermal patch does not cause any irritation and is safe for transdermal application [93].

Stability studies
The stability studies of optimised midazolam loaded niosome (N5) and midazolam loaded niosomal transdermal patch (NT5) was done in the refrigerator (4±2 °C) and room temperature (30±2 °C)/65±5%RH for 90 d[94]. The optimised midazolam loaded niosome was subjected to particle size analysis which reviewed that there are no major variations in its stability in both room and refrigerator temperatures as illustrated in fig. 9 (A) [95].

The entrapment efficiency of optimised midazolam loaded niosomal transdermal patch (NT5) was measured during different study intervals and illustrated in fig. 9 (B), confirmed that the formulations were stable at both the temperature conditions.

CONCLUSION
Epilepsy is characterized by the persistent increase of neuronal perturbation and inhibitory synaptic current breakdown which was treated by conventional antiepileptic drugs with high dosage and frequency of administration. The main purpose of this study was to develop a sustained delivery of midazolam loaded niosome through transdermal route for a patient who has frequent daily dosages where formulation effectively improved its release rate and permeation characteristics. Among the formulations prepared, midazolam loaded niosome (N5) and midazolam loaded niosomal transdermal patch (NT5) was fixed to be optimised formulations, which had maximum drug release of 97.3±0.35% and 98.9±0.20% and greater permeation than control drug solution. The flux and permeability coefficient was found to be higher for midazolam loaded niosomal transdermal patches when compared to control drug formulation which was statistically significant (P<0.05). The histopathological studies showed no significant alteration in its normal physiology when optimised formulations was permeated through porcine skin. The optimised formulations of niosome and transdermal patches were found be having good stability at room and refrigerator conditions. Thus formulated midazolam loaded niosomal transdermal patch represents to be an efficient and stable vesicular carrier for the transdermal delivery of an antiepileptic drug like midazolam which could be further studied by using in vivo animal models.
AUTHORS CONTRIBUTIONS
All the authors have contributed equally

CONFLICT OF INTERESTS
Declared none

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