Curcumin-loaded colloidal carrier system: formulation optimization, mechanistic insight, ex vivo and in vivo evaluation

Zrien Naz
Farhan Jalees Ahmad
Nanomedicine Research Lab, Faculty of Pharmacy, Jamia Hamdard, New Delhi, India

Abstract: The present work investigated the topical delivery potential of nanoemulsion gel loaded with curcumin (CR). CR nanoemulsion (CR-NE) was prepared by spontaneous emulsification method using oil (Labrafac PG/glyceryl triacetate), surfactant:cosurfactant (S:C) (tween 80/polyethylene glycol [PEG] 400) and water. The pseudo-ternary phase diagrams were constructed and thermodynamic stability testing was performed. Droplet size and zeta potential were evaluated using photon correlation spectroscopy and transmission electron spectroscopy. Six formulations selected with an average droplet size $\pm 70.2.72$ nm showed a fourfold increase in skin permeation as compared to crude CR solution in oil. The formulation CR-NE4 having a flux of $117.04 \pm 2.32 \mu g/cm^2/h$ and with maximum retention (42.87%) was selected, characterized (droplet size $=4.1.1.3 \pm 3.34$ nm and zeta potential $=-33.1 \pm 1.45$ mV), and incorporated into gel using carbopol-980 (1% w/v). Skin dynamics analyzed by confocal laser scanning microscopy showed maximum deposition of CR up to a depth of $86.98 \mu m$ and was in concordance with differential scanning calorimetry and Fourier transform infrared spectroscopy studies that confirmed lipid bilayer disruption, enhancing permeation. A 28-day anti-arthritis evaluation (body weight, paw edema, tibiotarsal joint thickness, TNF-$\alpha$ and IL-1$\beta$ levels, and histopathology) on Freund’s complete adjuvant induced arthritic rat model after topical application of CR-NE gel in Wistar rats demonstrated substantial reversal of arthritic symptoms. Thus, CR-NE gel possesses potential for therapeutic effects locally in inflammatory arthritic disorders with improved topical bioavailability.

Keywords: anti-arthritis, anti-inflammatory, curcumin, nanoemulsion, skin dynamics, topical

Introduction

Arthritic disorders are rheumatic diseases that are a major cause of locomotory dysfunction and disability in adult human population. The most prevalent forms of these articular diseases being osteoarthritis and rheumatoid arthritis. Rheumatoid arthritis is an auto immune disorder caused due to trigger of pro-inflammatory cytokines (eg, TNF-$\alpha$ and IL-1$\beta$) while osteoarthritis is a degenerative joint disease characterized by the progressive breakdown of articular cartilage. The pharmacotherapy for various forms of arthritis is achieved through symptomatic treatment with simple analgesics, steroidal and non-steroidal anti-inflammatory drugs, classically as the first line treatment followed by intra-articularly injected glucocorticoids and hyaluronic acid preparations as the disease progresses. The mainstay treatment, however has recurrent and potentially harmful side effects, low efficacy, and poor patient compliance. Several anti-inflammatory drugs have been withdrawn by the US Food and Drug Administration (FDA) due to severe cardiovascular side effects, and consequently,
plant-based agents are being promoted for the management of arthritis. Curcumin (CR), the yellow coloring agent found in turmeric (Curcuma longa, family Zingiberaceae), is a diferuloylmethane [1,7-Bis(4-hydroxy-3-methoxyphenyl)-1,6-heptadiene-3,5-dione] and has been used for centuries as a spice in the Indian subcontinent. It has been reported to have a wide range of activities like anti-inflammatory, anti-oxidant, anti-parasitic, anti-mutagenic, anti-cancer, and anti-arthritic (both in vitro and in vivo), as suggested by many studies. It has also been used in wound healing and as dietary supplement for arthritis in traditional medicine. The wide use of this drug, however, is limited by poor oral bioavailability due to extensive hepatic metabolism, low gastrointestinal absorption, and rapid elimination. Several nanoformulations of CR have been extensively investigated over the last decade like liposomes, nanoparticles, nanoconjugates, and nanoemulsions, which have been aimed at enhancing bioavailability by solubilization or encapsulation thus protecting against hydrolysis and achieving sustained release at the site of targeting. However, much attention has not been given to the topical formulation of CR which will aim at controlling local macrophage activity in synovial membrane by inhibiting the production of cytokines in arthritis related inflammation. Overcoming low oral bioavailability of CR is a big challenge, and it being highly lipophilic in nature (log P=2.92) shows good skin retention, therefore, topical delivery was intriguing. Nanoemulsion as a topical carrier system is very suitable with merits like low skin irritation, excellent permeation, and high drug-loading capacity as compared to other colloidal carriers. However, due to low retention at the site of application by virtue of low viscosity of nanoemulsion system and inconvenient application, it is often converted into a gel form.

Hence, in the present study a gel based colloidal carrier system loaded with CR intended to be used as adjunct therapy for the treatment of inflammation related to arthritic disorders was developed and investigated on a Freund’s complete adjuvant (FCA)-induced arthritic rat model. A mechanistic insight to the topical targeting potential of nanoemulsion gel (NEG) was investigated through skin dynamic study by differential scanning calorimetry (DSC), FT-IR microscopy and confocal laser scanning microscopy (CLSM) analysis of the treated rat skin.

**Materials and methods**

CR was purchased from Sigma Aldrich Co. (St Louis, MO, USA). Tween 80, and polyethylene glycol (PEG) 400 was procured from Hi-Media (Mumbai, India). Labrafac PG and Labrasol were generously gifted by Gattefosse, Bad Krozingen, Germany. Glyceroyl triacetate (triacetin) was purchased from Thomas Baker, Mumbai, India. High performance liquid chromatography (HPLC) grade methanol and acetonitrile were purchased from Merck Millipore, India. MilliQ water produced in-house by MilliQ water purification system (EMD Millipore, Billerica, MA, USA) was used for all the experiments. All other chemicals and reagents were of analytical grade.

**Solubility studies for excipient screening**

**Screening of oils**

In order to select the most suitable oil, solubility studies were conducted. An excess quantity of the drug (CR) was added to 1 mL of different oils (viz oleic acid, coconut oil, castor oil, sefsol, Labrafac PG, almond oil, isostearyl isostearate, olive oil, ethyl oleate, triacetin, arachis oil, sesame oil, and Miglyol 810n) taken separately in vials, until the loss of visual transparency. The vials were tightly stoppered and were continuously stirred for 24 hours at 37°C±0.5°C on a mechanical shaker. Samples were examined and the solubility was determined by the developed reversed-phased (RP)-HPLC analysis.

**Screening of surfactants and co-surfactants**

Similarly, for selection of the ideal surfactant tween 80, tween 60, tween 20, and Labrasol were tried. Likewise, PEG 400, Plurrol Oleique, propylene glycol, and Transcutol P were chosen as co-surfactants and drug solubility was determined as described under “Screening of oils”.

**Construction of pseudo-ternary phase diagrams**

Pseudo-ternary phase diagrams were constructed using aqueous titration method which includes stepwise addition of water to each weight ratio of oil and surfactants, and then mixing the components with the help of vortex mixer at 25°C. Surfactant:co-surfactant (S:C) ratios of 1:0, 1:1, 1:2, 1:3, 2:1, 3:1, and 4:1 were tried. Twelve different combinations in different weight ratios of oil and S:C, 1:9, 1:8, 1:7, 1:6, 1:5, 1:4, 1:3, 1:2, 1:1, 6:4 (1:0.7), 7:3 (1:0.43), and 9:1, were taken for each S:C ratio. For the determination of existence zone of nanoemulsion, the oil and S:C mixture was titrated with distilled water until a clear, isotropic and thermodynamically-stable dispersion with low viscosity was obtained. The volume of water was then recorded. The pseudo-ternary phase diagrams were constructed by...
plotting water phase, oil phase and $S_{\text{mix}}$ phase used in the experiment.

**Thermodynamic stability testing of nanoemulsion**

The best ratios from pseudo-ternary phase diagram having oil content 9%–12%, and $S_{\text{mix}}$ 50%–55% were selected and further subjected to thermodynamic stability evaluation. Parameters like opalescence, creaming and phase separation were observed.

**Centrifugation**

The formulations were centrifuged at 3,000 rpm at room temperature for 30 minutes and observed.

**Heating cooling cycle**

The formulations were stored at two different temperatures of 4°C–48°C for not less than 48 hours. Six cycles of this exposure were performed and stability was noted.

**Freeze-thaw cycle**

The same formulation was subjected to freeze-thaw cycle at temperature ranging from –21°C to 25°C for not less than 48 hours. The experiment was again performed six times and formulation was observed visually.

**Formulation of CR-loaded nanoemulsion**

CR-loaded nanoemulsions were prepared by first dissolving CR (0.25% w/v) in oil mix (Labrafac PG and glyceryl triacetate in equal ratio) and then slowly adding the required weight of $S_{\text{mix}}$ (tween 80 and PEG 400) and water, as calculated from the pseudo-ternary phase diagram, under constant magnetic stirring.

**Characterization of CR nanoemulsion (CR-NE)**

**Droplet size, size distribution, and zeta potential**

Droplet size and size distribution was determined using photon correlation spectroscopy that analyzes the fluctuations in light scattering due to Brownian motion of the droplets using Zetasizer Nano-ZS (Malvern Instruments, Malvern, UK). About 0.1 mL of the formulation was suitably diluted with water, mixed thoroughly with vigorous shaking and light scattering was monitored at 25°C at an angle of 90°. Zeta potential of the formulation was also determined using the same instrument.

**Transmission electron microscopy (TEM)**

TEM was performed to see the droplets’ morphology. A drop of the nanoemulsion was suitably diluted with distilled water (1:100), and applied on carbon-coated copper grid, negatively stained with 2% phosphotungstic acid and left for 30 seconds. Excess of phosphotungstic acid was removed by absorbing on a filter paper and the grid was analyzed using the Morgagni 268D transmission electron microscope (FEI Company, Hillsboro, OR, USA) operated at 60–80 kV at 1,550× magnification.

**RP-HPLC**

A simple RP-HPLC method was developed and validated for the estimation of CR in the bulk drug and nanoemulsion formulation. Shimadzu HPLC equipment comprising of quaternary LC-10AVP pumps, SPD-10AVP column oven, a variable-wavelength programmable UV-visible detector, and an-SCL-10AVP system controller consisting of 25 cm × 4.6 mm internal diameter (ID), 5-µm particle, 516 C18 DB reversed phase column (Supelco, Sigma-Aldrich, St Louis, MO, USA) was used. Mobile phase comprised of acetonitrile and 20 mM potassium dihydrogen orthophosphate (pH adjusted to 3.0) in the ratio 55:45 v/v, with a flow rate of 1 mL/min and injection volume of 20 µL. The detection wavelength was monitored at 420 nm. The concentration of drug was calculated by comparing the area peaks of unknown and standard samples with known concentrations.

**Preparation of skin samples**

Abdominal skin in full thickness was obtained from male Wistar rats weighing 224 ± 14.2 g from Institutional Animal Ethical Committee of Jamia Hamdard, New Delhi, India after approval from animal ethics committee. The hair on the skin surface was carefully shaved off using electrical razor, subcutaneous tissue was surgically removed and dermis side was wiped with isopropyl alcohol. The skin was washed with distilled water and visually inspected for integrity. A 3.0×3.0 cm² section of it was wrapped in aluminum foil and stored in a deep freezer at –20°C.

**Ex vivo skin permeation and deposition study**

Ex vivo permeation studies through rat skin were performed using an automated diffusion cell sampling system (SFDC6, Logan Instruments, Somerset, NJ, USA). Epidermal membrane samples were mounted onto the double-jacketed diffusion cells, 12 mL capacity and area 1.77 cm², equilibrated at 37°C±0.5°C for 24 hours. Amongst the nanoemulsion formulations that passed thermodynamic stability tests, six
formulations containing 0.25% w/v of CR each (CR-NE1, CR-NE2, CR-NE3, CR-NE4, CR-NE5, and CR-NE6), with particle size ≤70 nm and different S∞ ratios were selected and applied on the skin surface in the donor compartment. Aliquot (500 μL) was withdrawn from the receptor compartment at designated time intervals (viz 1, 2, 4, 6, 8, 10, 12, and 24 hours) for 24-hour period and analyzed for drug content by the developed RP-HPLC method. Receiver volume was immediately replenished with the same amount of medium-mixture of acetate buffer (pH 5.5) and ethanol in a ratio 7:3. Ethanol was added to the buffer to maintain the sink conditions. All the operations were carried out in triplicate. For skin deposition study, the skin was collected from the diffusion cell after the end of permeation study and homogenized in tissue homogenizer, the drug was extracted as per the developed liquid-liquid extraction method and analyzed. A control formulation (crude curcumin solution [C-CR]) containing CR (0.25% w/v) in the same oil was used for all comparisons.

Data analysis of skin permeation

The cumulative amount of drug permeated through the unit surface area corresponding to time, t, was calculated and plotted as a function of time. The steady state rate of skin permeation, flux (J, μg/cm²/h) was calculated from the slope of the permeation graph and the intercepts on the x-axis corresponded to the lag time. The permeability coefficient Kp was also calculated from the flux using the following equations:

\[ Qt = \frac{V_r C_r + \sum_{i=1}^{n} V_i C_i}{A} \]  
\[ K_p = \frac{J}{C_o} \]

where, Cr, Ci, and Co are the drug concentrations at time t at ith sampling and in the donor compartment, respectively. Vr and Vi is the volume of the receiver compartment and the sample, respectively.

Preparation of CR-NEG and CR crude gel (CR-CG)

The formulation showing the lower permeation flux was selected as the optimized one and converted into gel. In 10 mL of the CR-NE, 1% w/v Carbopol-980 was dispersed slowly with the help of a magnetic stirrer and kept in dark for 24 hours for complete swelling. The dispersion was neutralized using triethanolamine and propylene glycol was added as a plasticizer to obtain the gel. CR solution in ethanol (0.25% w/v) was prepared and incorporated in Carbopol-980 (1% w/v) gel (CR-CG) for comparative mechanistic and in vivo analysis.

Skin dynamics study

The freshly-prepared rat abdominal skin was washed with phosphate buffered saline (PBS) pH 7.4 thrice and mounted over a Franz diffusion cell with its epidermis facing the upper surface and 1 g of CR-NEG or CR-CG was placed in the donor compartment and sealed with parafilm. The receptor chamber was filled with acetate buffer (pH 5.5) and ethanol mixture (7:3) and left for 24 hours under constant magnetic stirring.

DSC study of skin

Approximately 5 mg of the skin was weighed and taken individually from the CR-NEG and CR-CG-treated skin, respectively, washed with PBS, blot-dried and hermetically sealed in aluminum pans for DSC studies. The scanning was done on a DSC6, differential scanning calorimeter (PerkinElmer Inc., Waltham, MA, USA) at a scanning rate of 10°C/min over the temperature range of 30°C–400°C. A DSC scan of each sample served as control for itself and the changes in the structures of both the skins were assessed and compared.

FT-IR microscopy study of the skin

The FT-IR spectra of the treated skin was recorded using Varian 7000 FT-IR spectrometer in conjunction with an UMA 600 microscope and a 128×128 focal-plane array detectors (FPA) detector, facility at Advanced Instrumentation Research Facility (AIRF), Jawaharlal Nehru University, New Delhi, India. The skin samples, treated separately with CR-NEG and CR-CG as described above, were taken for analysis. Later on, the skin samples were cut across the perimeter of the diffusion cell area and washed with PBS and air dried. Skin slides were prepared and mounted on the stage of microscope with coverslip removed and the FT-IR spectra of treated skin discs were recorded.

CLSM study

After 24 hours of permeation study, the skin was removed and washed with ethanol. The treated area was cut and mounted over glass slide with coverslip over it. CLSM was carried out using laser confocal microscope with fluorescence correlation spectroscopy, Olympus FluoView FV1000 (Olympus Corporation, Tokyo, Japan) with an argon laser beam at AIRF, Jawaharlal Nehru University, New Delhi. The fluorescent dye, Rhodamine B, being water-soluble, having an excitation wavelength of 520 nm, was incorporated in the CR-NEG to study drug deposition and penetration. CR,
itself being fluorescent with an excitation wavelength of 420 nm gave blue fluorescence along with Rhodamine B which gave red fluorescence.

**In vivo studies**

Animal studies were carried out after approval from Institutional Animal Ethics Committee, Jamia Hamdard, New Delhi (Protocol number 965) and guidelines laid down by Committee for the Purpose of Control and Supervision of Experiments on Animals were duly followed. A total of 30 male Wistar rats, having an average body weight 240±15 g, were randomly divided into five groups consisting of six animals each. Group I served as healthy normal control, Group II consisted of untreated arthritis-induced negative control, Groups III, IV, and V consisted of arthritis-induced animals treated with diclofenac-containing marketed gel, CR-NEG and CR-CG, respectively. The animals were housed in propylene cages in controlled conditions (temperature =20°C±2°C and humidity =55%±15%) and fed with standard pellet diet and water ad libitum.

**Dose calculation**

Dose calculation for herbal drugs is a difficult task due to lack of experimental data. However, in case of CR, acute oral toxicity in rats has been observed at a dose of >10,000 mg/kg, while its bioavailability is less than 25%. For topical application in rats, dose was calculated as follows. Effective human dose of CR (1,200 mg/kg) was divided by a conversion factor of 5/56 and the result was multiplied by its oral bioavailability.\(^{31}\) Thus, a topical dose of 25.71 mg/kg body weight was obtained.

**Disease induction and dosing schedule**

The animals were injected into the sub-planter region of the left hind paw, with 0.1 mL of FCA.\(^{32,33}\) This consisted of 5 mg *Mycobacterium butyricum* (Difco) being suspended in heavy paraffin oil (Merck Millipore) by thoroughly grinding with mortar and pestle to give a concentration of 5 mg/mL (0.05% w/v). This day was assigned as day “0”. Dosing with the drug formulation and with the standard was done twice on the same day and continued for 28 days, including the day of inoculation. A thick layer of the prepared gel was applied on the tibiotarsal joint in such a way as to cover the maximum area of the paw.

**Body weight measurement**

The body weight of all the animals in each group was measured on the day of initiation and the days specified (0, 4th, 7th, 14th, 21st, and 28th)\(^{34}\) and the change in body weight of each animal after the inoculation with FCA was calculated using the formula:

\[
\Delta BW(\%) = \left[ \frac{BW_i - BW_n}{BW_n} \right] \times 100
\]

where, \(\Delta BW(\%)\) is percent change in body weight in each group, \(BW_i\) is the average body weight of individual group on day of initiation and days specified, \(BW_n\) and is the average body weight of each group at the days specified.

**Tibiotarsal joint thickness (TJT) measurement**

The ankle (tibiotarsal) joint diameter of all the animals in each group was measured using a screw gauge on the specified days. Change in joint thickness was calculated by using the formula:\(^{34}\)

\[
TJT% = \frac{AD_i - AD_o}{AD_o} \times 100
\]

where, \(AD_i\) is the average ankle diameter of the groups on the days specified and \(AD_o\) is the average ankle diameter of the normal control group on that day.

**Hind paw edema volume (PV) measurement**

Paw volume of the left hind paw was measured plethysmographically on 0, 7th, 14th, and 28th day. The paw volume on the day “0” was taken as initial volume and difference in volume as an indicator of inflammation was measured for all the animals in all the groups on days given. Percent edema rate (ER \%) and arthritic index (AI\%) was calculated using the following formulae:

\[
ER\% = \frac{V_t - V_o}{V_o} \times 100
\]

\[
AI\% = \frac{E_t - E_o}{E_o} \times 100
\]

where, \(V_t\) is volume at specified day and \(V_o\) is the initial volume, \(E_t\) is the ER of the untreated control group, \(E_o\) is the ER of the test groups. The area under the curve of increase in paw volume and time gives the rate of inhibition of arthritis (In\%) by different treatment. It can also be calculated as per the following equation:

\[
In\% = 100 - \left[ \frac{(ER_t - ER_o)}{ER_o} \right] \times 100
\]
where \( ER_t \) is the average ER of the treated groups, \( ER_c \) is the average ER of the normal control and \( ER_u \) is the average ER of the untreated control group.

**Preparation of joint tissue homogenate**

At the end of the experiment, the paw at the level of malleolus was amputated and the digits were trimmed off. The skin and underlying tissue at the joint were carefully removed using a surgical blade. An approximate area of the tibiotarsal joint was taken and carefully dipped in liquid nitrogen. The joint was grounded and homogenized using a tissue homogenizer (Remi Instruments, Mumbai, India) using 10× volume of PBS at pH 7.4 at 15,000 rpm for 30 minutes. This was later centrifuged and the supernatant was collected for the biochemical estimation of TNF-\( \alpha \) and IL-1\( \beta \).

**Biochemical estimation of cytokines in serum**

Blood was collected in a sterile centrifuge tube from the tail vein of all groups using a micro-capillary tube on the 28th day and left undisturbed at 37\(^\circ\)C for 45 minutes. During this period, the serum exuded and the clot retracted. The serum was aspirated using a sterile pipette after centrifugation at 3,000 rpm for 15 minutes. The serum samples obtained by centrifugation at room temperature were used for the biochemical estimation of TNF-\( \alpha \) and IL-1\( \beta \) using ELISA (enzyme-linked immunosorbent assay) kit supplied by RayBio Lab, New Delhi, India and manufactured by RayBiotech, Inc., Norcross, GA, USA.

**Histopathological examination**

At termination of the study, the tibiotarsal joint was transected at the level of the medial and lateral malleolus and was collected into formalin for histopathological evaluation. Ankle joints were collected into 10% neutral buffered formalin for at least 24 hours prior to placement in Surgipath Decalcifier I (Leica Microsystems, Wetzlar, Germany) for approximately 1 week. After decalcification, the digits were trimmed and the ankle joint was transected in the longitudinal plane to give approximately equal halves. Joints were processed for paraffin embedding, sectioned, and stained with hematoxylin and eosin for cartilage changes.

**Statistical analysis**

All the quantitative data were expressed as an average ± SD for each group. Comparisons were made using the Dunnett’s \( t \)-test. In all cases \( P<0.05 \) were considered statistically significant.

**Results and discussion**

**Solubility studies for excipient screening**

In the present study, different oils, natural as well as semi-synthetic, were chosen and solubility of CR was determined. The selection of the oil phase was made in such a way that a maximum amount of CR should get incorporated and it should be best suited for topical application. Therefore, the oil phase comprising of Labrafac PG and glyceryl triacetate (triacetin) in equal ratio was selected. Glyceryl triacetate, being an excellent solvent blended well with the oil and enhanced the solubility of CR.

For the selection of surfactant and co-surfactant solubilization efficiency was taken into account. The surfactant chosen must be able to lower the interfacial tension to a very small value, provide a flexible film that can readily deform around droplets, and have the appropriate hydrophilic lipophilic balance (HLB) value to provide the correct curvature at the interfacial region for the desired nanoemulsion type. Transient negative interfacial tension and fluid interfacial film is rarely achieved by the use of single surfactant, usually necessitating the addition of a co-surfactant.

Therefore, tween 80, a non-ionic, non-irritant surfactant having an HLB value of 15 was selected as the surfactant and PEG 400 as the co-surfactant.

**Construction of pseudo-ternary phase diagrams and formulation of CR-NE**

The pseudo-ternary phase diagrams were constructed for different ratios of tween 80 and PEG 400. The surfactant and co-surfactant mass ratios greatly affected the area of nanoemulsion formation on the phase diagram. Higher concentration of co-surfactant \( S_{mix} \) (1:2, 1:3, 1:4) led to narrower zone of nanoemulsion region with higher frequency of breakdown of nanoemulsion upon storage, which can be attributed to the fact that increase in co-surfactant concentration might lead to destabilization effect. A wider zone of nanoemulsion was obtained from \( S_{mix} \) ratios in the order of 2:1 < 4:1 > 3:1 (Figure 1A). This phenomenon has been supported by the fact that increase in surfactant concentration leads to lowering of the interfacial barrier and formation of thermodynamically-stable nanoemulsion owing to reduction in size, high surface area and charge.

Further optimization of the nanoemulsion was carried out by the stability test comprising of heating and cooling cycles, centrifugation and freeze-thaw cycle. Different
formulations from pseudo-ternary phase diagram of each SMix ratio were selected based on the criteria that they should possess an optimum oil concentration ranging from 9% to 12% and surfactant and co-surfactant ratio of 50%–55%. The SMix ratios of 1:0, 1:1, and 1:2 showed turbidity or phase separation upon thermodynamic stability testing. This may be due to increase in particle diameter upon centrifugation. Opalescence is observed which may be reversed on standing. Heating results in randomness due to increase in enthalpy, while cooling results in loss of entropy or coalescence, leading to an increase in particle size. 37

Characterization of optimized formulation

The evaluation of morphology by TEM revealed that the size of the nanoemulsion was less than 50 nm (Figure 1B), which was in concordance with the photon correlation spectroscopy analysis. The average hydrodynamic particle size and poly dispersity index (PDI) of the optimized formulation (CR-NE4) were found to be 41.13±3.34 nm and 0.173±0.04, respectively (Figure 1C). The zeta potential of the formulation was found to be −33.1±1.45 mV, as evident from Figure 1D. The size of the nanoemulsion globule less than 50 nm, having a dispersibility of less than one is considered ideal for topical delivery, as it would provide a large surface area. 38,39 The nanoemulsion globules were spherical with a smooth flexible boundary. The negative zeta potential value of 30 mV or less was considered sufficient to prevent coalescence and maintain the interfacial boundary of the nanoemulsion. 25

Ex vivo permeation and deposition study

Six formulations, two each from SMix ratios of 2:1, 3:1, and 4:1 having an oil content (Labrafac PG: glyceryl triacetate =1:1) 9%-12% v/v and SMix range 50%-55% v/v, were selected and characterized for droplet size (Figure 2A). CR at a concentration of 0.25% w/v was mixed thoroughly with the oil phase. Then, the required amount of SMix and water was added under constant magnetic stirring.
The comparative permeability study was performed for the selected nanoemulsions CR-NE1, CR-NE2, CR-NE3, CR-NE4, CR-NE5, and CR-NE6 through the excised rat skin. The steady state flux and permeability coefficient were compared with that of control, C-CR (Table 1). The selected formulations owing to their low particle size (Z-average ≤70 nm), greatly enhance the rate of permeation. The S<sub>mix</sub> blend increases the solubilization efficiency and permeability coefficient of the skin as the interfacial barrier decreases and partitioning improves owing to increase in lipophilicity.<sup>40</sup> It is evident that the steady state flux (J<sub>ss</sub>) and the permeability coefficient of CR-NE4 are 117.03 ± 2.32 µg/cm<sup>2</sup>/h and 4.68 × 10<sup>-3</sup> cm/h, respectively, which are the lowest amongst the nanoemulsions, but higher than the CR solution in oil, C-CR (Figure 2B). Enhancement ratio showed that there was a fourfold increase in the steady state flux in case of nanoemulsion, with P-value < 0.0001, which is highly significant (Table 1). In topical drug delivery, a low rate of permeation and high retention is desirable. The percent skin deposition of the drug in the six investigated nanoformulations were

Table 1  Permeation data of selected curcumin nanoemulsions and curcumin solution in oil

<table>
<thead>
<tr>
<th>Serial no</th>
<th>Code</th>
<th>Z-average (nm)</th>
<th>J&lt;sub&gt;ss&lt;/sub&gt; (µg/cm&lt;sup&gt;2&lt;/sup&gt;/h)</th>
<th>K&lt;sub&gt;p&lt;/sub&gt;×10&lt;sup&gt;-3&lt;/sup&gt;</th>
<th>ER</th>
<th>% DD</th>
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<tr>
<td>1</td>
<td>CR-NE1</td>
<td>57.99±2.41</td>
<td>149.45±4.02&lt;sup&gt;a&lt;/sup&gt;</td>
<td>5.97</td>
<td>4.07±1.36</td>
<td>28.47±1.42</td>
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<td>2</td>
<td>CR-NE2</td>
<td>62.38±1.38</td>
<td>153.54±8.61&lt;sup&gt;a&lt;/sup&gt;</td>
<td>6.14</td>
<td>4.18±2.60</td>
<td>22.68±2.32</td>
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<td>3</td>
<td>CR-NE3</td>
<td>44.29±1.64</td>
<td>127.50±5.44&lt;sup&gt;a&lt;/sup&gt;</td>
<td>5.09</td>
<td>3.47±1.80</td>
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<td>4</td>
<td>CR-NE4</td>
<td>41.13±3.34</td>
<td>117.04±2.32&lt;sup&gt;a&lt;/sup&gt;</td>
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<td>3.18±1.43</td>
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<td>50.75±2.56</td>
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<td>58.77±2.45</td>
<td>159.79±6.57&lt;sup&gt;a&lt;/sup&gt;</td>
<td>6.39</td>
<td>4.35±1.87</td>
<td>21.00±2.62</td>
</tr>
<tr>
<td>7</td>
<td>C-CR</td>
<td>–</td>
<td>36.71±2.98</td>
<td>1.46</td>
<td>1.00</td>
<td>9.12±1.23</td>
</tr>
</tbody>
</table>

Notes: <sup>a</sup>Skin flux at steady state; <sup>b</sup>permeability coefficient. Z-average: mean hydrodynamic droplet size; value in mean ± standard deviation, n=3. <sup>c</sup>P-value < 0.0001 (mean flux of CR-NE1–6 versus C-CR).

Abbreviations: CR-NE, curcumin nanoemulsion; C-CR, crude curcumin solution in oil; DD, drug deposition; ER, enhancement ratio.
found to be in the range of 20%–43% with CR-NE4 showing the highest skin retention of 42.87%±1.72% which is significantly higher (P<0.05) than the C-CR which showed only 9.12%±1.23% drug retention. Thus, CR-NE4 was selected as the optimized formulation for further studies.

Preparation of NEG

The formulation CR-NE4 was selected as the optimized one on the basis of its permeation data and was converted into gel by using carbopol-980 (1% w/v) (Table 2) for mechanistic and in vivo investigations. The gel matrix further lowers the permeation rate and allows free movement of globules within due to hydration resulting in increase in the surface area.41 It provides ease of application and holds drug topically on affected area for a longer duration.

Skin dynamics study

DSC study of skin

DSC study of the skin provides a direct measurement of the changes in the skin surface based on enthalpy changes. The mechanism of permeation enhancement by the NEG could be elucidated by the thermograms obtained from the CR-NEG treated skin, while the one treated with CR-CG served as the control. The DSC thermograms of the CR-CG skin showed the presence of peaks Pa and Pb obtained at 80°C–110°C, respectively, which can be attributed to melting of stratum corneum lipids (Figure 3A). In the DSC thermograms of CR-NEG treated skin the peaks at Pa and Pb shifted further lower by 10°C–20° indicating enhanced fluidization of the lipids. In addition to this, an additional peak Pc appeared at temperature 142°C which may be due to incorporation of the nanoemulsion into stratum corneum.

The FT-IR microscopy study of the skin

After the 24 hours permeation study, the skins were removed from the Franz diffusion cell and the treated area was excised, analyzed, and compared for the characteristic FT-IR bands. There is a pronounced difference in the FT-IR spectra of the CR-NEG treated skin when compared with the CR-CG treated one which served as control (Figure 3B). Characteristic peaks in the FT-IR spectra of the CR-CG treated skin appeared at 2,920 cm⁻¹ and 2,849 cm⁻¹, which corresponds to asymmetric and symmetric C-H stretching, respectively. Amide I band C=O stretching appeared near 1,650 cm⁻¹. The C-H stretching wave numbers are related to lipid order in the skin and C=O is related to protein conformation.43 The peak intensities were high and predominantly mimicked the spectra of untreated skin with slight shift in wavenumber which can be attributed to hydration of the skin.43 The peak height decreased in CR-NEG-treated skin for C-H stretching which can be contributed to the lipid bilayer conformational disruption and fluidization of membrane lipids. The nanoemulsion caused a change in fluidity of lipids, and therefore the characteristic peaks of C-H stretching at 2,850 cm⁻¹ and 2,920 cm⁻¹ were of weaker intensity.

CLSM studies

CLSM images of the Rhodamine B dye-loaded CR-NEG were compared with that of C-CR (Figure 4). The figure shows intense red and blue fluorescence corresponding to Rhodamine B and CR, respectively. The intensity of red fluorescence increases consistently with the depth of skin up to 58.98 μm and faded afterward, while blue fluorescence was highest at the depth of 86.98 μm (Figure 4A–D). This could be attributed to the fact that oil-in-water (o/w) nanoemulsion created a diffusion barrier as a result of partitioning across lipid bilayer of the stratum corneum. Rhodamine B being hydrophilic followed the polar pathway of permeation, while CR diffused from the inner core following the continuous non-polar pathway.44 Skin integrity has been altered by the nanoemulsion proving that nanoemulsion has the potential to permeate across the stratum corneum. The drug got released due to the increase in concentration gradient created as a result of partitioning following Fickian diffusion.45 The lipophilicity increased with the thickness of the skin, hence blue fluorescence increased gradually indicating higher concentration of CR.

In vivo studies

Body weight measurement

Body weight measurement data after FCA injection showed a close relationship between the extent of joint inflammation

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Table 2 Composition of different curcumin formulations

<table>
<thead>
<tr>
<th>Ingredients (% w/v)</th>
<th>C-CR</th>
<th>CR-NE</th>
<th>CR-CG</th>
<th>CR-NEG</th>
</tr>
</thead>
<tbody>
<tr>
<td>Curcumin</td>
<td>0.25</td>
<td>0.25</td>
<td>0.25</td>
<td>0.25</td>
</tr>
<tr>
<td>Carbopol-980</td>
<td>–</td>
<td>–</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>Labrafac PG</td>
<td>6.0</td>
<td>6.0</td>
<td>–</td>
<td>6.0</td>
</tr>
<tr>
<td>Glyceryl triacetate</td>
<td>6.0</td>
<td>6.0</td>
<td>–</td>
<td>6.0</td>
</tr>
<tr>
<td>Tween 80</td>
<td>–</td>
<td>–</td>
<td>44*</td>
<td>44*</td>
</tr>
<tr>
<td>PEG 400</td>
<td>–</td>
<td>11*</td>
<td>–</td>
<td>11*</td>
</tr>
<tr>
<td>TEA</td>
<td>–</td>
<td>0.05</td>
<td>–</td>
<td>0.05</td>
</tr>
<tr>
<td>Water (qs)</td>
<td>–</td>
<td>100</td>
<td>100</td>
<td>100</td>
</tr>
</tbody>
</table>

Note: *±8, 4:1.
Abbreviations: TEA, triethanolamine; PEG, polyethylene glycol; C-CR, crude curcumin; CR-NE, curcumin nanoemulsion; CR-CG, curcumin crude gel; CR-NEG, curcumin nanoemulsion gel; S, surfactant; cosurfactant; qs, quantity sufficient.
and the degree of body weight loss (Figure 5A). Significant weight loss was observed between days 14, 21, and 28 in group II ($P<0.001$). However, groups receiving treatment with CR-NEG and standard drug did not show any significant weight loss after day 14 owing to the effect of treatment. Moreover, the shielding effect of CR-NEG was significantly greater as compared to standard formulation after the inflammatory phase subsided after day 14.

Figure 3 Skin dynamics study by DSC and FT-IR.
Notes: (A) DSC thermograms and; (B) FT-IR peaks of skin treated with CR-NEG and CR-CG, respectively.
Abbreviations: FT-IR, Fourier transform infrared spectroscopy; CR-NEG, curcumin nanoemulsion gel; CR-CG, curcumin crude gel; DSC, differential scanning calorimetry.
Figure 4 CLSM images of the rat skin treated with CR-NEG and CR-CG showing permeation of Rhodamine B (red fluorescence) and curcumin (blue fluorescence) at different depths of skin.

Notes: (A, B) Red and blue fluorescence at a depth of 86.98 µm; (C) combined fluorescence at depth of 86.98 µm; (D) maximum red fluorescence at a depth of 58.98 µm; (E) combined fluorescence in different skin layers after 24 hours.

Abbreviations: CLSM, confocal laser scanning microscopy; CR-NEG, curcumin nanoemulsion gel; CR-CG, curcumin crude gel.

TJT measurement
Upon injection with FCA, there was inflammation at the site and joint at day 4, there was noticeable difference in %TJT in groups II, III, IV, and V, as compared to the normal control. Significant enhancement was found in group II on 4, 7, and 14 days compared to healthy control (Figure 5B). After day 14 until 28 the rate of inflammation decreased. Reversal was observed in all the treatment groups with varying magnitude. Treatment with CR-NEG was found to be almost as effective as standard in controlling tibiotarsal joint inflammation.

PV measurement
Between 0 and 7th day insignificant differences in PV were found among all the groups. Ominously increasing trend was seen in PV for group II from day 1 onward until day 4, when there is maximum inflammation (Figure 5C). Afterward, inflammation started declining until day 14 but there was reversal and inflammation could be noticed on the non-injected paws as well owing to secondary lesions and all the comparisons were made between 4th, 14th, 21st, and 28th days’ observations. Mean AIs of the groups and %In in different treatment groups are shown in Table 3. AI is used to define the severity of joint inflammation based on calculation of scores of the paws after arthritis induction as compared to the normal control. The AIs of CR-NEG treated group on day 4 and 14 were 49.67%±6.42% and 198.20%±7.91%, respectively, and %In on day 14 and 28 were 70.50%±8.17% and 85.30%±1.72%, respectively, which were significantly higher than both standard (P<0.05) and CR-CG (P<0.01) treatments. The days 4–14 mark the proliferative phase which is characteristic of inflammation, hyperplasia and macrophage activities in the synovial membrane. The rate of inhibition was clearly better in case of CR-NEG as compared to CR-CG (P<0.01).
Biochemical estimation of cytokines in serum and joint tissue homogenate

The pro-inflammatory cytokines, TNF-α and IL-1β were significantly enhanced in blood as well as joint tissue on 28th day of induction of arthritis in group II rats as compared to group I (healthy control) (P<0.01). TNF-α and IL-1β concentrations were markedly reduced in group III, IV, and V. However, most significant effect was seen in CR-NEG treated group followed by standard treatment group (Figure 5D, E). This trend was observed in both the serum as well as in joint tissue homogenate. But, at the level of control the effect was more pronounced in tissue homogenate as compared to serum mainly due to etiology of arthritic disorders. TNF-α is secreted by macrophages, monocytes, neutrophils, T-cells and NK-cells following their stimulation by bacterial lipopolysaccharide. TNF-α shares many of the pro-inflammatory activities of IL-1β, and may therefore also promote synovitis and stimulate cartilage and bone breakdown.

The affected joints synthesize and release IL-1β, IL-6, and TNF-α which are synergistic in inflammatory activity and TNF-α may be an essential stimulus for the production of other cytokines such as IL-1β in the arthritic joint since neutralizing antibodies directed against TNF-α reduce the spontaneous production of IL-1β from synovial mononuclear cells derived from the joint. Thus, the elevated TNF-α and IL-1β levels in group II indicated the extent of cartilage damage. The CR-NEG showed most significant depression of both TNF-α and IL-1β levels as compared to CR-CG which can be attributed to high permeability achieved through nanoemulsion.
Table 3: Arthritic index (AI%) and rate of inhibition of arthritis (In%) calculated from hind paw volume for each group

<table>
<thead>
<tr>
<th>Groups</th>
<th>Day 4</th>
<th>Day 7</th>
<th>Day 14</th>
<th>Day 21</th>
<th>Day 28</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>%AI ± SD</td>
<td>%In ± SD</td>
<td>%AI ± SD</td>
<td>%In ± SD</td>
<td>%AI ± SD</td>
</tr>
<tr>
<td>I*</td>
<td>141.66 ± 16.41</td>
<td>-</td>
<td>1023.65 ± 4.24</td>
<td>-</td>
<td>2374.19 ± 14.78</td>
</tr>
<tr>
<td>II</td>
<td>0 ± 0</td>
<td>-</td>
<td>0 ± 0</td>
<td>-</td>
<td>0 ± 0</td>
</tr>
<tr>
<td>III</td>
<td>42.66 ± 4.29</td>
<td>30.60 ± 4.86</td>
<td>96.77 ± 4.19</td>
<td>50.14 ± 8.32</td>
<td>143.62 ± 5.72</td>
</tr>
<tr>
<td>IV</td>
<td>49.67 ± 6.42</td>
<td>33.89 ± 0.02</td>
<td>128.18 ± 3.40</td>
<td>57.14 ± 159</td>
<td>198.20 ± 7.91</td>
</tr>
<tr>
<td>V</td>
<td>19.74 ± 6.94</td>
<td>17.18 ± 8.03</td>
<td>73.41 ± 5.50</td>
<td>43.30 ± 43</td>
<td>86.37 ± 9.93</td>
</tr>
</tbody>
</table>

Notes: *Arthritis index of normal animal was compared with control; *rate of inhibition in normal and control was insignificant; **level of significance (P<0.01); *** (P<0.05) as compared to standard (group III).

Abbreviation: SD, standard deviation.

In the present study, an NEG loaded with CR was developed and characterized for topical administration in the management and treatment of arthritis. The skin dynamical studies revealed that topical treatment with CR could effectively reduce the inflammatory response, as evidenced by a reduction in the arthritic index and rate of inhibition of arthritis. The nanocarrier system provided an insight into the mechanism of topical delivery of drugs like CR. The in vivo permeation and deposition studies clearly indicated that nanocarrier system as the delivery system has high potential for drug delivery to the target site. The DSC thermograms and the FT-IR scans of the CR-NEG treated skin provided an insight into the potential of topical delivery of drugs like CR.

Conclusion

In the present study, CR-loaded NEG was better in controlling the progression of arthritis than the standard marketed formulation upon topical application. Thus, from the above findings, it can be concluded that CR-loaded NEG is effective in suppressing the arthritic symptoms. Therefore, this nanoemulsion-based curcumin-loaded colloidal carrier system has better potential in the management and control of arthritis progression.

Histopathological examination

The representative photomicrographs of histological sections are shown in Figure 6. Group I (Figure 6A), being normal control revealed smooth synovial epithelium with normal joint cavity without any sign of cellular infiltration or bone erosion. Group II (Figure 6B) rats showed marked cellular infiltration and synovial hyperplasia, joint cavity, cartilage destruction, and bone erosion. Group III (Figure 6C), being control marketed gel, showed marked cellular infiltration and bone erosion. However, CR-CG (Figure 6D) did not seem to have comparable effects as that of groups III and IV. The histological sections confirmed the increased effects on cellular infiltration, joint space narrowing, and synovial hyperplasia, but markedly reduced the process of cartilage destruction and bone erosion. Group IV (Figure 6E) showed marked cellular infiltration and synovial hyperplasia, joint cavity, cartilage destruction, and bone erosion as compared to healthy control. The representative photomicrographs of histological sections were shown in Figure 6. The above findings confirmed that CR-loaded NEG was better in controlling the progression of arthritis than the standard marketed formulation upon topical application.
References


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Disclosure

The research work is original and the authors report no conflict of interest.

Notes:

1. Figures 6, A–E. Photomicrographs of histological section of ankle joint in the respective groups. 
2. Group I: normal joint; Group II: control; Group III: treatment with standard marketed gel; Group IV: treatment with CR-CG; and Group V: treatment with CR-NEG. Images taken on 10× magnification.

Abbreviations: CR-NEG, curcumin nanoemulsion gel; CR-CG, curcumin crude gel.


