Effects of nanoparticles with hydrotropic nicotinamide on tacrolimus: permeability through psoriatic skin and antipsoriatic and antiproliferative activities

Abstract: The hybrid system based on nanoparticles (NPs) self-assembled by the conjugations of hyaluronic acid with cholesterol (HA–Chol NPs) combined with nicotinamide (NIC) for tacrolimus (FK506), ie, FK506 NPs–NIC, has been confirmed to exhibit a significant synergistic effect on FK506 permeation through and into intact skin; thus, it may be a promising approach for FK506 to effectively treat skin diseases. The aim of this study was to evaluate its potential for the treatment of psoriasis. In vitro permeation through the psoriatic skin was carried out, and the results revealed that the combination of NPs with NIC exhibited a significant synergistic effect on FK506 deposition within the psoriatic skin (3.40±0.67 μg/cm²) and penetration through the psoriatic skin (30.86±9.66 μg/cm²). The antipsoriatic activity of FK506 NPs–NIC was evaluated through the treatment for imiquimod (IMQ)-induced psoriasis. The psoriasis area and severity index (PASI) score demonstrated that FK506 HA–Chol NPs–NIC exerted the effect on ameliorating the skin lesions comparable to clobetasol propionate (a positive drug for psoriasis) and superior to commercial FK506 ointment (Protopic®), and the histological study showed that it presented a synergistic effect on antipsoriasis after FK506 incorporation into NPs combined with NIC hydrotropic system, which might ultimately increase the therapeutic effect and minimize the systemic side effects by reducing the overall dose of FK506. RAW 264.7 cell uptake presented the enhancement of drugs delivered into cells by HA–Chol NPs–NIC. The antiproliferative activity on HaCaT cells identified that FK506 HA–Chol NPs–NIC exhibited significant inhibiting effects on HaCaT proliferation. The results support that the combination of HA–Chol NPs with NIC is a promising approach for FK506 for the treatment of psoriasis.

Keywords: tacrolimus, nanoparticles, nicotinamide, percutaneous delivery, psoriasis, antiproliferation

Abbreviations
ANOVA, analysis of variance; C6, coumarin 6; Chol, cholesterol; CLSM, confocal laser scanning microscope; CP, clobetasol propionate; CCTCC, China Center for Type Culture Collection; DAPI, 4′,6-diamidino-2-phenylindole; DMEM, Dulbecco’s Modified Eagle’s Medium; DMSO, dimethyl sulfoxide; EDC, ethylcarbodiimide hydrochloride; FBS, fetal bovine serum; FDA, Food and Drug Administration; FK506, tacrolimus; HA, hyaluronic acid; HaCaT, immortalized human keratinocytes; IL-2, interleukin-2; HE, hematoxylin and eosin; HPLC, high-performance liquid chromatography; IMQ, imiquimod; MTT, 3-(4,5-dimethyl-2-thiazolyl)-2,5-diphenyl-2-H-tetrazolium bromide; LSD, least significant difference; NF-κB, nuclear factor-κB;
NHS, N-hydroxysuccinimide; NS, normal saline; OD, optical density; PASI, psoriasis area and severity index; SC, stratum corneum; SD, standard deviation; NIC, nicotinamide; NPs, nanoparticles; PARP-1, poly(ADP-ribose) polymerase-1; PBS, phosphate buffer solution.

Introduction

Psoriasis is one of the most common chronic immune-mediated inflammatory erythematousquamous dermatoses, prevalent among 2%–5% of the world population. It is characterized by keratinocyte hyperproliferation, hyperplasia of the epidermis, and massive infiltration of inflammatory leukocytes in the epidermis and dermis.1–3 The clinical feature of visible disfiguration of erythematous skin lesions covered with white silvery scales at the skin surface often inflicts upon patients through psychological burdens and decreased quality of life.

Currently, the treatment of psoriasis composes of a wide range of treatment options, including topical agents (corticosteroids and vitamin D analogs), systemic administration (cyclosporine A, methotrexate, and retinoids), phototherapy (psoralen combination with ultraviolet radiation A [PUVA] and ultraviolet radiation B [UVB]), and biological treatment (specific siRNA).4–6 Up to date, topical therapy remains a preferred treatment strategy for psoriasis because the site of action is direct. The challenge for topical treatment of psoriasis is the poor drug penetration through the psoriatic skin owing to the physiopathology of psoriasis. Psoriatic skin with the characteristics of thickening and abnormal maturation of the SC, lipid imbalance, and deficiency of natural moisturizers hinders drug penetration and limits therapeutic efficacy.9 A novel and improved percutaneous delivery system to overcome the SC barrier of psoriatic skin is vital to antipsoriatic drug for topical therapy.

Psoriasis often requires long-term and routine administration of anti-inflammatory drugs such as corticosteroids. Topical application of corticosteroids is the first-line therapies for psoriasis; however, its utility is likely to induce skin atrophy by inhibition of collagen synthesis and increased susceptibility to skin infection.10 Since the inflammation in psoriasis is related to an exacerbated immune response, immunosuppressant drug is a valid therapeutic strategy. FK506, a macrolide immunosuppressive drug isolated from Streptomyces tsukubaensis, has been proved to be useful in the treatment of psoriasis and psoriasis arthritis when administered systemically.11–14 Its mechanism is similar to that of cyclosporine A, a potent cyclic undecapeptide immunosuppressant, which is therapeutically effective in patients with severe psoriasis.15 Moreover, FK506 is ~10–100-fold better than cyclosporine A in inhibiting the mass production of lymphokines involved in IL-2 production, and therefore it has been widely used in organ transplantations.16 FK506 achieves immunosuppression by the inhibition of IL-2 and other immune response cytokine transfection, thereby preventing T-lymphocyte activation,17–19 while activated T cells are necessary for the development and persistence of psoriasis lesions. Clinical research data have indicated that systemic administration of FK506 can play an important role in the treatment of severe and recalcitrant psoriasis on the face, genitalia, intertriginous areas, and corporal plaques.20 FK506 is available commercially as an ointment formulation (Protopic®) approved by the FDA for topical treatment of atopic dermatitis. In clinical practice, FK506 ointment is commonly used off-label in other skin diseases, including psoriasis, and FK506 was recommended in particular for the treatment of facial and intertriginous psoriasis by the American Academy of Dermatology.21 However, topical application of the commercial ointment failed to promote FK506 deposition in hyperkeratotic psoriatic plaques.22 Meanwhile, FK506 ointment is difficult to be washed off from the skin surface and may create a sticky sense and discomfort to the patient due to its greasy nature. Several reports demonstrated that topical application of FK506 ointment was only effective under occlusion or to some specific regions of the body where the skin was thinner and more permeable, such as the face and intertriginous areas.23–25 The poor efficacy of FK506 for topical therapy was attributed to its high hydrophobicity and high molecular weight (822.05 Da), which prevent it from penetrating into and through the skin. In our previous study,26 we have formulated a hybrid nanoparticle system for FK506 percutaneous delivery. The system was based on the combination of amphiphilic conjugations of HA–Chol-self-assembled NPs and hydrotropic NIC, ie, FK506 HA–Chol NPs–NIC, and it has been found that either NP carrier or NIC hydro tropy could enhance the permeation of FK506 through the intact skin, and the combination of HA–Chol NPs with NIC could synergistically enhance the permeation of FK506 into deeper epidermal and dermal layers of the skin in comparison with commercial FK506 ointment. These results supported that the hybrid system facilitated FK506 in the treatment of skin diseases.

In this study, the aim is to investigate the potential of FK506 HA–Chol NPs–NIC for the treatment of psoriasis. FK506 HA–Chol NPs–NIC was formulated by self-assembling. The murine psoriasis model was constructed by topical application of IMQ on mouse dorsal skin. In vitro permeability through the psoriatic skin of FK506 from HA–Chol NPs–NIC was
evaluated in comparison with HA–Chol NPs, NIC complex solution, and FK506 ointment. The antipsoriatic activity of FK506 HA–Chol NPs–NIC against IMQ-induced psoriasis was assessed in comparison with FK506 NPs, FK506 NIC complex solution, FK506 ointment, and CP (a positive reference), and the effect was evaluated with the PASI score and histological examination. Finally, RAW 264.7 cells were used to assess the cellular uptake using a hydrophobic probe of C6 and HaCaT cells were used to evaluate the antiproliferative activity of FK506 HA–Chol NPs–NIC.

Materials and methods

Materials

Cholesterol chloroformate, N-(3-dimethylaminopropyl)-N′-EDC, NHS, and pyrene were purchased from Aladdin (Shanghai, China). Sodium hyaluronate (HA, Mw 30 kDa) was obtained from Bloomage Freda Biopharm Co., Ltd (Shandong, China). Anhydrous ethylenediamine was purchased from Damao Reagent (Tianjin, China). Trimethylamine was obtained from Zhiyuan Reagent (Tianjin, China). NIC was a gift of Guangzhou Changlong (Guangzhou, China). FK506 was purchased from Teva Czech Industries s.r.o. (Opava-Komarov, Czech Republic). FK506 ointment (Protopic®) was obtained from Astellas Toyama Co., Ltd. (Toyama, Japan). The product 502 super glue was purchased from Henggu Plastic Industry Co., Ltd (Zhejiang, China). Anhydrous ethylenediamine was purchased from Sigma-Aldrich (St Louis, MO, USA). FBS and DMEM were purchased from Gibco (Gaithersburg, MD, USA). Other reagents were of analytical grade.

Murine RAW 264.7 macrophage cells were obtained from the Experimental Animal Center of Sun Yat-sen University (Guangzhou, China). HaCaT cells were obtained from CCTCC (Wuhan, China).

Preparation and characterization of FK506 NPs

FK506 NPs were prepared as described in our previous study. In brief, amphiphilic HA–Chol conjugates were synthesized by amide bond formation of the –NH₂ of cholesteryl amine and the –COOH of HA. The purified HA–Chol conjugates and FK506 were dispensed in 20% (w/v) NIC solution or aqueous solution with the final FK506 concentration of 1 mg/mL, and then, FK506-loaded HA–Chol NPs with NIC or without NIC were obtained by sonicating for 6 min using ultrasonic cell disruptor (JY92-2D; Ningbo Xinzi Science and Technology Co., Ningbo, People’s Republic of China). The particle size and zeta potential of FK506 NPs–NIC and FK506 NPs were determined by Nano ZS Zetasizer (Malvern Instruments, Malvern, UK), and they were ~210 nm and ~42 mV and ~240 nm and ~43 mV, respectively. The encapsulation efficiency of FK506 NPs–NIC and FK506 NPs was ~79 and 72%, respectively.

Psoriasis mouse model induced by IMQ

BALB/c mice (male, 6–8 weeks old) were purchased from the Experimental Animal Center of Sun Yat-sen University. The mice were housed under specific pathogen-free conditions at constant levels of humidity and temperature on 12 h light/dark cycles, provided with food and water ad libitum. All procedures performed using animals were in accordance with the Principles of Laboratory Animal Care and Use in Research published by the Chinese Ministry of Health, and the protocols were approved by the Institutional Animal Care and Use Committee of Sun Yat-sen University.

The dermatitis induced by topical application of IMQ on mouse dorsal skin closely resembles human plaque-type psoriasis lesions not only with respect to phenotypic presentation and histological characteristics but also in the development of the inflamed scaly skin lesions. Therefore, the murine psoriasis model was constructed in this study by topical application of IMQ on mouse dorsal skin. In brief, the mice were administrated a daily topical dose of 5 mg IMQ with a commercially available cream (Hubei Keyi Pharmaceutical Co., Ltd) on the shaved dorsal skin for 10 consecutive days. The treated area was examined daily for skin inflammation or lesions, and the clinical PASI was scored based on the extent of erythema, scaling, and epidermal thickening of the inflammation on the dorsal region. After 10 days, the mice were sacrificed by cervical dislocation under sodium pentobarbital anesthesia and the treated dorsal skin was immediately excised. Punch biopsies of lesioned skin samples were fixed in 4% paraformaldehyde and paraffin embedded for histological analysis to confirm the psoriasis-like skin inflammation induced by IMQ.

In vitro permeation through psoriatic skin

The experiment was conducted with Franz diffusion cell (TK-12A; Kaikai Technology Co., Ltd, Shanghai, China) under occlusive conditions with an effective contact area of 3.14 cm² and a receptor volume of 7.5 mL. After the psoriasis was successfully induced by IMQ as described in the
“Psoriasis mouse model induced by IMQ” section, the mice were sacrificed by cervical dislocation under sodium pentobarbital anesthesia. The dorsal skin with psoriatic lesions was immediately excised, and the subcutaneous tissues were surgically removed; then, the treated psoriasis skin was sandwiched between the donor and receptor cells with SC facing upward into the donor. The receptor medium was NS containing 25% ethanol with respect to the sink condition for the permeation of FK506. Skin temperature was maintained at 32°C±1°C throughout the experiment with a circulating water bath and stirred at 250 rpm. After equilibration, 500 μL of FK506 HA–Chol NPs–NIC, HA–Chol NPs, and FK506 NIC complex suspension and 0.5 g of commercial preparation of FK506 ointment were applied onto the skin. In our preliminary experiment, FK506 in the receptor medium could not be detected until 10 h permeation in the groups with NIC and 12 h permeation in the group without NIC; thus, the receptor medium at intervals of 12, 14, 16, 18, 20, and 24 h after permeation were sampled, and equal volume of fresh receptor medium was replenished after each sampling.

After 24 h permeation, the diffusion cells were dismantled, the residual formulations on the treated skin were carefully removed, and the skin was cut into pieces. Two volume of methanol was added to the skin pieces for extracting FK506 retained in the psoriatic skin. The extraction was performed by sonicating for 30 min following centrifuging for 5 min at 5,000 rpm. The supernatant was collected for the samples.

All the samples from the above were filtered with 0.22 μm microporous nylon membrane and analyzed by HPLC. The HPLC system (1,260 series; Agilent Technologies, Santa Clara, CA, USA) consisted of quaternary pump (G1310A), degasser (G1322A), autosampler (G1329), column thermostat (CO-1000), ultraviolet (UV) detector (G1314A), and data processing software (Agilent Chem Station for LC systems). A C8 column (Grace®, 250×4.6 mm, 5 μm) was used for FK506 analysis with water/isopropyl alcohol/tetrahydrofuran (5:2:2, v/v/v) as the mobile phase at a flow rate of 0.8 mL/min at 55°C, and the detection wavelength was set at 220 nm.

**Antipsoriatic activity assessment**

**Treatment with different formulations of FK506**

The animals were randomly divided into nine groups (nine mice per group) as follows: control group, IMQ group, IMQ–vehicle group, IMQ–NIC group, IMQ–FK506 ointment group, IMQ–FK506–NIC group, IMQ–FK506 NPs group, IMQ–FK506 NPs–NIC, and IMQ–CP group. Except that the control group (disease-free control group) was not treated with IMQ, each mouse of other eight groups was treated with IMQ as described in the “Psoriasis mouse model induced by IMQ” section to induce psoriasis, and then the animals were treated by applying different formulations onto the IMQ-induced dorsal region for 10 consecutive days. However, IMQ group was the disease model group without the treatment of other drugs. IMQ–vehicle group was treated with the vehicle containing excipients in the delivery system without FK506. IMQ–NIC group was treated with 20% NIC solution, which was investigated to confirm whether NIC ameliorated the psoriasis symptom or not. IMQ–FK506 ointment group, IMQ–FK506–NIC group, IMQ–FK506 NPs group, and IMQ–FK506 NPs–NIC group were treated with different formulations containing 0.1% FK506 (equal to the content of FK506 in commercial ointment), and the different formulations were topically administered onto the IMQ-induced psoriasis lesions at an equal dose of 20 μg/cm² of FK506 in each group. CP, as an extremely potent synthetic steroid hormone approved for the management of psoriasis by FDA,30 was used as a positive control drug to exhibit ameliorating symptoms in this study, and it was topically administered once daily at a dose of 8 μg/cm² in the IMQ–CP group. All the therapy lasted for 10 days.

After 10 days’ treatment, mice were sacrificed by cervical dislocation under sodium pentobarbital anesthesia and the treated dorsal skin was immediately excised. Punch biopsies of lesional skin samples from each mouse in each group were fixed in 4% paraformaldehyde and paraffin embedded for histological analysis.

**Scoring severity of skin inflammation**

To monitor and grade the severity of the psoriasis-like skin condition on days 0, 2, 4, 6, 8, and 10, the scoring system was based on the clinical PASI5,29,31,32 Three parameters, including erythema, scaling, and thickening, were assigned independently on a scale from 0 to 4 as follows: 0, none; 1, slight; 2, moderate; 3, marked; 4, very marked. A scoring table with red tints was used to score the level of erythema. The cumulative score (erythema, scaling, and thickening) denoted the severity of inflammation (scale 0–12). After 10 days treatment, mice were sacrificed by cervical dislocation under sodium pentobarbital anesthesia. In addition of evaluation by PASI, the treated dorsal skin was immediately excised, and the thickness of the skinfold was measured in sextuplicate using a digital micrometer with an accuracy of ±0.02 mm (Mitutoyo Corporation, Mitsutomo, Japan).

**Histopathological analysis**

The paraffin-embedded punch biopsies were sectioned longitudinally into 6 μm thick sections. Then, the tissue
sections were HE stained for histological and microarchitecture evaluation. Blinded to treatment group, epidermal thickness, an accepted end point for measuring psoriasis severity, was measured as an average of 20 random measurements of the distance from the SC to the deepest part of the epidermis. Experiments were conducted in sextuplicate, and the data were averaged to evaluate epidermal proliferation and inflammation.

Cell culture

The cells were cultured in DMEM supplemented with FBS (10%, v/v), 100 IU/mL penicillin, and 100 μg/mL streptomycin in the humidified atmosphere of 95% air with 5% CO₂ at 37°C (Forma 3111; Thermo Fisher Scientific, Waltham, MA, USA). The medium was replenished every other day, until the monolayer cultures reached 60%–80% confluency. Subsequently, the culture medium was aspirated off and the cells were then washed with PBS, followed by incubation with trypsin-EDTA solution (1 mL) at 37°C for 5 min. The trypsin was deactivated by the addition of 2 mL of DMEM supplemented with FBS. Then, the cell suspension was centrifuged at 1,000 rpm for 3 min, and the resulting supernatant was removed. Eventually, 4 mL of fresh culture medium was added to resuspend the detached cells. The suspension was then ready to be used for experimental work.

Cellular uptake of fluorescence-labeled formulations

RAW 264.7 cells were cultured in 15 mm² confocal dish (1×10⁵ cells/well) with DMEM. After cultured for 24 h, the culture medium was removed and washed with PBS for three times. Two hundred microliter of HA–Chol NPs–NIC, HA–Chol NPs, NIC complex suspension, and aqueous suspension containing equal amount of C6 (0.1%, w/v) were diluted by DMEM without FBS to 10 μg/mL and added to the dish. After culturing for 4 h, the culture medium was removed and the cells were washed with PBS for three times. Then, the cells were fixed with 4% paraformaldehyde solution at ambient temperature for 15 min, and the nucleus of cell was labeled by DAPI. RAW 264.7 cells without adding fluorophore were tested as a blank control group. The cellular uptake was evaluated using CLSM (LSM 710; Zeiss, Oberkochen, Germany). The excitation and emission wavelengths of C6 were 430 and 485 nm, respectively.

MTT cell proliferation assay

Immortalized human HaCaT keratinocytes were used as the model of proliferating epidermal cells to test the antiproliferative efficacy of different formulations containing FK506. 33–36 HaCaT cells in the log phase were seeded into 96-well plates (Corning Inc., Corning, NY, USA) with a density of 5×10³ cells/well and maintained in an atmosphere of 5% CO₂ at 37°C for 24 h. HA–Chol NPs–NIC, HA–Chol NPs, NIC complex suspension, and aqueous suspension containing equal amount of FK506 (0.1%, w/v) were diluted by culture medium to the final concentration at 10 μg/mL of FK506, and empty NPs–NIC vehicle was diluted as well to be the vehicle control. These different preparations were added to the cells cultured in 96-well microplates. Cell proliferation was assessed at different time points (6 and 18 h). The cells with no formulations were designated as normal control cells, while blank control cells contained only culture medium were used for colorimetric zero adjustment. The medium was then removed, and the adhering cells were washed with PBS for three times. The culture fluid was then replaced with fresh DMEM without FBS containing 0.5 mg/mL of MTT. After incubation for 4 h at 37°C, the MTT reaction medium was removed from the plates and 150 μL of DMSO was added to solubilize formazan blue. The plates were shaken for 15 min in the dark to dissolve the formazan completely. The OD was then recorded at 490 nm with an ELISA Microplate Reader (Synergy H1; BioTek, Winooski, VT, USA). All tests were performed in triplicate, and results were calculated as a percentage of the values of the test wells over the corresponding normal control wells.

Data analysis

Each experiment was performed at least in triplicate. The data were given as mean ± SD. The statistical differences between the groups used one-way ANOVA followed by LSD as post hoc analysis (SPSS Version 19.0; SPSS Inc., Chicago, IL, USA). P<0.05 was considered statistically significant.

Results and discussion

Permeability through psoriatic skin

The proliferation and abnormal differentiation of keratinocytes of psoriasis render the psoriatic skin with the thickening of the SC and abnormal maturation of the SC, which exhibits abnormal epidermal barrier. Permeability through psoriatic skin comes closer to the clinical application than that through intact skin, which facilitates to elucidate the exact influence of drug delivery system on antipsoriatic drugs percutaneous delivery. Therefore, the psoriatic skin was used as the skin barrier to perform permeation study. The cumulative permeated amount of FK506 through psoriatic skin was plotted versus time and the retention of FK506 in the skin, as shown in Figure 1. FK506 ointment (0.1%, w/w) was taken as the reference group because there was no significant difference
between FK506 ointment and 0.1% FK506 suspension in our preliminary experiment. As shown in Figure 1, during the 24 h permeation, the FK506 permeation from NIC complex or HA–Chol NPs was significantly higher than that from FK506 ointment (P<0.05) throughout the experiment. After 24 h permeation, the cumulative permeated amount of FK506 from NPs or NIC complex was 2.38-fold or 1.71-fold as that of FK506 ointment, and the skin retention of FK506 from NPs or NIC complex was 2.54-fold or 1.96-fold as that of FK506 ointment. Furthermore, FK506 from NPs–NIC showed the highest permeation at all time points in comparison with NPs or NIC complex group, and the cumulative permeated amount of FK506 was 30.86±9.66 μg/cm² after 24 h permeation, which was significantly higher than that of NIC complex with 14.21±4.15 μg/cm² (P<0.05) or NPs alone with 19.72±8.06 μg/cm² (P<0.05), and the FK506 retention in the skin from NPs–NIC presented the highest retention of 3.40±0.67 μg/cm² than NPs of 2.34±0.35 μg/cm² or NIC complex of 1.80±0.59 μg/cm² (P<0.05). The results indicated that either NPs carrier or NIC hydrotropy could enhance the permeation of FK506 through and into the psoriatic skin, and the combination of HA–Chol NPs with NIC could synergistically enhance the permeation of FK506 into and through the psoriatic skin. The mechanism of these systems enhancing drug permeation through psoriatic skin might be attributed to the following facts. For NPs, the encapsulating FK506 accumulated on the skin surface to create high local concentration of FK506 and then facilitated the drug’s sustained release and permeation through the skin. For NIC complex, FK506 permeation was probably facilitated in two ways. First, psoriatic skin presented drier or more rigid than the intact skin,40 and NIC exhibited moisturizing effect on the skin,39 which would modify the barrier property and enhance the penetration of drugs. Second, the hydrotropic effect of NIC on FK506 increased its solubility and provided FK506 concentration gradient and partition of skin/vehicle. Taken together, the hybrid system of HA–Chol NPs combined with NIC took both advantages of NPs and NIC to enhance FK506 permeation through and retention in the psoriatic skin. The combination of HA–Chol NPs with NIC might be a promising approach of FK506 for topical antipsoriatic treatment.

**Effects of FK506 HA–Chol NPs–NIC on ameliorating the incassation and inflammation of the psoriatic skin**

The skin disorders of IMQ-induced psoriasis include erythema, scales, and thickness. Two or three days after the initiation of IMQ application, the treated dorsal skin of mouse started to display typical symptoms of erythema, scaling, and thickening. The clinical PASI was scored based on the extent of erythema, scales, and thickness on days 0, 2, 4, 6, 8, and 10, and the average scores were calculated. The score

![Figure 1](https://www.dovepress.com/)

**Figure 1** Permeation profiles of FK506 through the psoriatic skin of mouse from different formulations in vitro (A) and FK506 skin retention after 24 h permeation of different formulations in vitro (B).

**Notes:** Each symbol and bar represent the mean ± SD of six determinations. Significant differences were calculated using the ANOVA test (*P<0.05). The concentration of FK506 in each formulation was 1 mg/mL.

**Abbreviations:** ANOVA, analysis of variance; FK506, tacrolimus; NIC, nicotinamide; NPs, nanoparticles; SD, standard deviation.
of erythema, scales, or thickness was plotted as a function of time (Figure 2A–C, respectively). The total score was calculated as the sum of erythema, scales, and thickness, and it was also plotted as a function of time (Figure 2D). At the time point of day 8, representative clinical features of the mouse were photographed (Figure 3A), and the thickness of the skinfold was measured (Figure 3B). As shown in Figure 2, the severity increased with continuous application of IMQ and the PASI score peaked on day 8. After treating with different formulations containing 0.1% FK506, the symptoms were alleviated with different levels. Reduction in the PASI score indicated the degree of healing of the treatment. The PASI score of IMQ–FK506–NIC group, IMQ–FK506 NPs group, or IMQ–FK506 NPs–NIC group was significantly less than that of IMQ group (P<0.05), which was comparable to IMQ–CP (P>0.05) throughout the experiment.

Figure 2 The severity of each psoriatic symptom; erythema (A), scaling (B), skin thickening (C), and total score (D) over time.

Notes: The severity of each psoriatic symptom was evaluated with the PASI score of psoriatic dorsal regions of mouse. Scoring was performed on days 0, 2, 4, 6, 8, and 10, based on the erythema, scales, and thickness of PASI; each symbol represents the mean ± SD of nine animals. Each symptom was assigned a score of 0–4 as follows: 0, none; 1, slight; 2, moderate; 3, marked; and 4, very marked. The total score was the sum of erythema, scaling, skin thickening. (×: control group, □: IMQ group, +: IMQ–vehicle group, *: IMQ–NIC group, ○: IMQ–Protopic® group, ●: IMQ–FK506–NIC group, ■: IMQ–FK506 NPs group, ▲: IMQ–FK506 NPs–NIC group, and ▽: IMQ–CP group).

Abbreviations: CP, clobetasol propionate; FK506, tacrolimus; IMQ, imiquimod; NIC, nicotinamide; NPs, nanoparticles; PASI, psoriasis area severity index.
or FK506–NIC and FK506 NPs–NIC. Second, FK506 with 0.1% concentration after NIC solubilizing or NPs encapsulating had obtained the maximum therapeutic effect comparable to that of CP; thus, the combination of NPs with NIC might reduce the overall dose of FK506 for antipsoriasis. However, the hypothesis will be the subject to further research.

IMQ–NIC and IMQ–vehicle groups presented slight reduction in the score and a little improvement in the appearance of dorsal skin, which might be attributed to the effect of NIC. In the cosmetic industry, NIC was widely used in topical preparations due to its moisturizing and depigmenting properties. The existing preclinical data show that NIC has the potential for topical application in cosmetic or pharmaceutical industry because of its barrier restoration, anti-inflammatory, and depigmenting effects.\textsuperscript{41,42} Moreover, NIC has a therapeutic potential as an inhibitor of proinflammatory cytokines in inflammatory disease.\textsuperscript{43} Activation of NF-κB is a very important factor in both lesional and nonlesional skin from patients with psoriasis. NIC was an inhibitor of PARP-1, which regulated the expression of inflammatory cytokines, adhesion molecules, chemokines, and inflammatory mediators (such as interleukin-12, interleukin-23, and tumor necrosis factor-α) through enhancing NF-κB-mediated transcription. This mechanism implied that suppression of NF-κB was involved in NIC-resulted anti-inflammation, antiproliferative, and antipsoriatic effects.\textsuperscript{43,44} It was also reported that NIC may prove a useful addition to the antipsoriatic repertoire and may enhance the efficacy of calcipotriene when used in combination for topical psoriasis treatment.\textsuperscript{45} Therefore, NIC might play a dual role of hydrotrope for water-insoluble FK506 and adjuvant therapeutic agent in the hybrid system of FK506 NPs–NIC for antipsoriasis.

After 10 days’ treatment, the thickness of the skinfold from the IMQ–FK506–NIC group, IMQ–FK506 NPs group, or IMQ–FK506 NPs–NIC group was thinner than that from

\begin{figure}[h]
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\caption{Representative photography of mouse after 8 days treatment with different formulations (A) and skinfold thickness measurement (B). \textbf{Notes:} Each bar represents the mean ± SD of nine determinations. Significant differences were calculated using the ANOVA test (*\textit{P} < 0.05). The concentration of FK506 in each formulation was 1 mg/mL. \textbf{Abbreviations:} ANOVA, analysis of variance; CP, clobetasol propionate; FK506, tacrolimus; IMQ, imiquimod; NIC, nicotinamide; NPs, nanoparticles; SD, standard deviation.}
\end{figure}
the IMQ group, and comparable to that of IMQ–CP, while there was no significant difference between IMQ–NIC, IMQ–vehicle, or IMQ–Protopic® and IMQ groups. Taken together, FK506–NIC, FK506 NPs, or FK506 NPs–NIC can significantly ameliorate the visible inflammation signs of IMQ-induced psoriasis.

**Histological analysis**

The HE staining of dorsal skin was performed to observe the histological changes of the skin induced with IMQ and then treated with different formulations. As shown in Figure 4, when the dorsal skin was induced by IMQ, it presented an increased epidermal hyperplasia, elongation of rete-like ridges, acanthosis in the epidermis, and perivascular infiltration of the inflammatory cells in the upper dermis, resembling closely a phenotype typical of human psoriatic skin. Additionally, abnormal keratinocyte differentiation with marked hyperkeratosis was observed. The hyperplasia of basal and suprabasal keratinocytes resulted in epidermal thickening. The epidermal thickness of each group was measured in microscope (Figure 5). The epidermal thickness of the control group without any treatment was 16.1±4.24 μm, while the epidermal thickness after IMQ-induced was 88.7±15.90 μm, and that of IMQ–NIC, IMQ–vehicle, or IMQ–Protopic® treatment was 73.5±8.70, 69.6±10.36, or 57.1±12.51 μm, respectively. IMQ–FK506 ointment presented a reduction in epidermal thickness, which confirmed that FK506 was effective in antipsoriasis; however, the effect needed to be improved. The epidermal thickness was significantly reduced in the IMQ–FK506–NIC group with 25.9±6.34 μm (P<0.05), IMQ–FK506 NPs group with 26.4±6.09 μm (P<0.05), IMQ–FK506 NPs–NIC group with 20.6±7.13 μm (P<0.05), and IMQ–CP group with 21.7±6.32 μm (P<0.05). The reduction of effect on epidermal thickening was in the order of FK506 NPs–NIC = CP > FK506–NIC = FK506 NPs. Compared to FK506–NIC or FK506 NPs, FK506 NPs–NIC presented a synergistic effect on antipsoriasis and comparable to that of CP. The histological study provided further evidence that FK506 NPs–NIC ameliorated IMQ-induced psoriasis and presented a synergistic effect on antipsoriasis after FK506 incorporation into NPs combined with NIC hydro tropic system.

Based on the antipsoriatic activity, HA–Chol NPs–NIC was supposed to reduce the overall dose and drug consumption of FK506, by effectively solubilizing, encapsulating, and localizing at the applied target site and will ultimately increase the therapeutic effect and minimize the systemic side effects.

![Figure 4](https://www.dovepress.com/)

**Figure 4** Histological analysis of skin after treatment with different formulations.

**Notes:** The epidermal thickness was used to assess the epidermal hyperplasia. The mean of epidermal thickness was calculated based on 20 random site measurements.

**Abbreviations:** ANOVA, analysis of variance; CP, clobetasol propionate; FK506, tacrolimus; IMQ, imiquimod; NIC, nicotinamide; NPs, nanoparticles.
uptake by RAW 264.7 indicated that the system of HA–Chol NPs with NIC facilitated to deliver the drug to immune cell and to exert an effect on immune-mediated skin disease.

**Antiproliferative effect of different formulations**

Psoriasis was mainly characterized by an aberrant hyperproliferation of keratinocyte in the interfollicular epidermis. Inhibition of epidermal cell proliferation is very important in the treatment of psoriasis. To evaluate the inhibiting effect of different formulations on HaCaT cell proliferation, cell viability was measured with time. As shown in Figure 7, after incubation with different formulations for 6 h, no cytotoxicity was observed in the cells treated with NPs–NIC vehicle. Both FK506–NIC and FK506 NPs–NIC significantly delayed the growth of HaCaT in comparison with the control group indicating the antiproliferative effects on HaCaT, while the other groups containing the equal amount of FK506 (0.1%) such as FK506 suspension and FK506 NPs presented no significant inhibiting effects on HaCaT. Based on these comparisons, the inhibiting effects of FK506–NIC and FK506 NPs–NIC on HaCaT growth under 6 h incubation are mainly attributed to the solubilization of NIC on FK506, which ensured the efficient concentration of FK506. While FK506 was encapsulated in NPs, it released in a sustained manner from NPs; therefore, FK506 was under the efficient concentration. However, after 18 h incubation, the formulations including FK506–NIC, FK506 NPs, FK506 NPs–NIC, and FK506 suspension exhibited significantly inhibiting effects on HaCaT growth with the cell viability of 53%±5%, 83%±6%, 61%±7%, and 83%±13%, respectively. Combined the results from 6 and 18 h, it can be concluded that FK506 exerts the inhibiting effect on HaCaT growth in a time-dependent manner. Interestingly, NPs–NIC vehicle also inhibited the growth after incubation for 18 h, and the cell viability of FK506–NIC group decreased more than that of the other group. These results may be attributed to the high solubility of FK506 with the NIC hydrotropy and that NIC itself can promote the antiproliferative effect. Indeed, our preliminary studies showed that NIC markedly impaired the proliferation and differentiation of HaCaT cell line. As expected, HA–Chol NPs–NIC showed a strong inhibitory effect on the proliferation of HaCaT cells. However, there was no significant difference between the HA–Chol NPs–NIC group and the FK506–NIC group. Overall, these results show that FK506 HA–Chol NPs–NIC could efficiently inhibit the proliferation of HaCaT and have a promising prospect in the treatment of psoriasis.

**Cellular uptake of C6-loaded NPs into RAW 264.7**

Psoriasis is an immune-mediated inflammatory skin disease; since macrophages and dendritic cells play key roles in the immune response, delivery of FK506 to these cells would facilitate the immunotherapy. Therefore, cellular uptake of drugs by murine macrophages was investigated. We chose RAW 264.7 cell to evaluate the cellular uptake of HA–Chol NPs. C6 was used as the hydrophobic fluorescence probe to mimic the hydrophobic drug, and the cellular uptake was observed by CLSM. The uptake of C6 by RAW 264.7 cell from HA–Chol NPs–NIC, HA–Chol NPs, 20% (w/v) NIC solution, and aqueous suspension containing equal amount of C6 (10 μg/mL) was evaluated (Figure 6). As shown in Figure 6A, RAW 264.7 cells presented the nucleus labeled by DAPI in the DMEM group without background green fluorescence interference. After 4 h incubation, Figure 6B presents negligible fluorescence in the C6 suspension group indicating very little C6 uptake by RAW 264.7 cells, and Figure 6C shows that NIC solution could slightly enhance the cell uptake of C6 compared to that of C6 suspension. However, when C6 was loaded in HA–Chol NPs, the fluorescence intensity was significantly strengthened, indicating that its uptake by RAW 264.7 cells was significantly enhanced (Figure 6D). When C6 was loaded in HA–Chol NPs with NIC, the fluorescence intensity was a little stronger than that of HA–Chol NPs alone (Figure 6E). The results of the cellular uptake by RAW 264.7 indicated that the system of HA–Chol NPs alone (Figure 6F). The results of the cellular NIC, the fluorescence intensity was a little stronger than that (Figure 6D). When C6 was loaded in HA–Chol NPs with its uptake by RAW 264.7 cells was significantly enhanced fluorescence intensity was significantly strengthened, indicating that the cell uptake of C6 compared to that of C6 suspension.

**Notes:** The epidermal thickness measured under the microscope. The mean of epidermal thickness was calculated based on 20 random site measurements. Significant differences were calculated using the ANOVA test (P<0.05). The concentration of FK506 in each formulation was 1 mg/mL.

**Abbreviations:** ANOVA, analysis of variance; CP, clobetasol propionate; FK506, tacrolimus; IMQ, imiquimod; NIC, nicotinamide; NPs, nanoparticles.

**Figure 5** The epidermal thickness measured under the microscope.

**Figure 6** The fluorescence intensity was a little stronger than that of HA–Chol NPs alone (Figure 6F). The results of the cellular NIC, the fluorescence intensity was a little stronger than that (Figure 6D). When C6 was loaded in HA–Chol NPs with its uptake by RAW 264.7 cells was significantly enhanced fluorescence intensity was significantly strengthened, indicating that the cell uptake of C6 compared to that of C6 suspension.

**Psoriasis was mainly characterized by an aberrant hyperproliferation of keratinocyte in the interfollicular epidermis.** Inhibition of epidermal cell proliferation is very important in the treatment of psoriasis. To evaluate the inhibiting effect of different formulations on HaCaT cell proliferation, cell viability was measured with time. As shown in Figure 7, after incubation with different formulations for 6 h, no cytotoxicity was observed in the cells treated with NPs–NIC vehicle. Both FK506–NIC and FK506 NPs–NIC significantly delayed the growth of HaCaT in comparison with the control group indicating the antiproliferative effects on HaCaT, while the other groups containing the equal amount of FK506 (0.1%) such as FK506 suspension and FK506 NPs presented no significant inhibiting effects on HaCaT. Based on these comparisons, the inhibiting effects of FK506–NIC and FK506 NPs–NIC on HaCaT growth under 6 h incubation are mainly attributed to the solubilization of NIC on FK506, which ensured the efficient concentration of FK506. While FK506 was encapsulated in NPs, it released in a sustained manner from NPs; therefore, FK506 was under the efficient concentration. However, after 18 h incubation, the formulations including FK506–NIC, FK506 NPs, FK506 NPs–NIC, and FK506 suspension exhibited significantly inhibiting effects on HaCaT growth with the cell viability of 53%±5%, 83%±6%, 61%±7%, and 83%±13%, respectively. Combined the results from 6 and 18 h, it can be concluded that FK506 exerts the inhibiting effect on HaCaT growth in a time-dependent manner. Interestingly, NPs–NIC vehicle also inhibited the growth after incubation for 18 h, and the cell viability of FK506–NIC group decreased more than that of the other group. These results may be attributed to the high solubility of FK506 with the NIC hydrotropy and that NIC itself can promote the antiproliferative effect. Indeed, our preliminary studies showed that NIC markedly impaired the proliferation and differentiation of HaCaT cell line. As expected, HA–Chol NPs–NIC showed a strong inhibitory effect on the proliferation of HaCaT cells. However, there was no significant difference between the HA–Chol NPs–NIC group and the FK506–NIC group. Overall, these results show that FK506 HA–Chol NPs–NIC could efficiently inhibit the proliferation of HaCaT and have a promising prospect in the treatment of psoriasis.
Conclusion

The hybrid system of HA–Chol NPs combination with NIC for FK506 has been confirmed to exhibit a significant synergistic effect on FK506 permeation through and into the intact skin; therefore, it may be a promising approach for FK506 to effectively treat skin diseases. In this study, in vitro permeation through the psoriatic skin indicated that either NIC or HA–Chol NPs alone significantly enhanced FK506
permeation through and into the psoriatic skin, and the combination of HA–Chol NPs with NIC exhibited a significant synergistic effect on the permeation in comparison with NIC or HA–Chol NPs alone. The treatment for IMQ-induced psoriasis demonstrated that FK506 HA–Chol NPs–NIC had an ameliorating effect on the skin lesions comparable to that of CP and superior to that of FK506 ointment and that it presented a synergistic effect on antipsoriasis after FK506 incorporation into NPs combined with NIC hydrotropic system, which might ultimately increase the therapeutic effect and minimize systemic side effects by reducing the overall dose and consumption of FK506. RAW 264.7 cell uptake presented the enhancement of drugs delivered into cells by HA–Chol NPs–NIC. The antiproliferative activity on HaCaT cells identified that FK506 HA–Chol NPs–NIC had an ameliorating effect on HaCaT prolifera
tion. In conclusion, the results indicate that the combination of HA–Chol NPs with NIC is a promising approach for FK506 for the treatment of psoriasis. Nevertheless, further works on the reduction of FK506 dose and the immune-related molecular mechanism of antipsoriasis need to be carried out to clarify the potential of HA–Chol NPs–NIC for FK506 as a percutaneous delivery system for the treatment of psoriasis.

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Disclosure
The authors report no conflicts of interest in this work.

References


