Transportan in nanocarriers improves skin localization and antitumor activity of paclitaxel

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Abstract: In this study, the ability of nanocarriers containing protein transduction domains (PTDs) of various classes to improve cutaneous paclitaxel delivery and efficacy in skin tumor models was evaluated. Microemulsions (MEs) were prepared by mixing a surfactant blend (polyoxyethylene 10 oleoyl ether, ethanol and propylene glycol), monocaprylin, and water. The PTD transportan (ME-T), penetratin (ME-P), or TAT (ME-TAT) was added at a concentration of 1 mM to the plain ME. All MEs displayed nanometric size (32.3–40.7 nm) and slight positive zeta potential (+4.1 mV to +6.8 mV). Skin penetration of paclitaxel from the MEs was assessed for 1–12 hours using porcine skin and Franz diffusion cells. Among the PTD-containing formulations, paclitaxel skin (stratum corneum + epidermis and dermis) penetration at 12 hours was maximized with ME-T, whereas ME-TAT provided the lowest penetration (1.6-fold less). This is consistent with the stronger ability of ME-T to increase transdermal water loss (2.4-fold compared to water) and tissue permeability. The influence of PTD addition on the ME irritation potential was assessed by measuring interleukin-1α expression and viability of bioengineered skin equivalents. A 1.5- to 1.8-fold increase in interleukin-1α expression was induced by ME-T compared to the other formulations, but this effect was less pronounced (5.8-fold) than that mediated by the moderate irritant Triton. Because ME-T maximized paclitaxel cutaneous localization while being safer than Triton, its efficacy was assessed against basal cell carcinoma cells and a bioengineered three-dimensional melanoma model. Paclitaxel-containing ME-T reduced cells and tissue viability by twofold compared to drug solutions, suggesting the potential clinical usefulness of the formulation for the treatment of cutaneous tumors.

Keywords: microemulsion, nanocarriers, protein transduction domains, paclitaxel, skin, transdermal

Introduction

It is well known that drug delivery into/across the skin is restricted by the cutaneous barrier provided mainly (but not exclusively) by the stratum corneum (SC).1 Drugs displaying molecular weight <500 g/mol and log P between 0.5 and 2.5 are considered good candidates for topical delivery.2 However, the majority of the drugs available do not meet these requirements, and penetration-enhancing strategies are needed to improve their skin penetration. One example is the antitumor agent paclitaxel; as a large and lipophilic drug (MW = 853.9 g/mol; log P =3.5), its penetration is limited.3 Only formulations for systemic use are currently available, but despite their efficacy against tumors affecting the skin, severe systemic adverse effects limit their use in dermatology.4,6 Thus, nanocarriers capable of improving paclitaxel cutaneous localization while limiting its transdermal permeation could potentially allow the use of this drug for the treatment of skin tumors.3,5,7,8 In this context, this study was aimed at evaluating the cutaneous delivery, safety, and efficacy of...
paclitaxel nanocarriers in bioengineered models of healthy and tumor-containing skin.

Different types of nanocarriers have been investigated for topical paclitaxel delivery; however, cutaneous localization of the drug is not often given priority over transdermal delivery during formulation design. Thus, in the first part of this study, the ability of a selected nanocarrier to improve the amount of drug delivered into skin layers was optimized. Microemulsions (MEs) were selected as nanocarriers due to their simple preparation (since there is no special requirement in terms of energy supply), thermodynamic stability, and versatility of composition, which can reduce costs. Considering that drug penetration can be enhanced when cationic lipids, protein transduction domains (PTDs), and pore-forming antimicrobial peptides are included in nanocarriers, we evaluated the effects of adding three different types of PTDs on the microemulsion safety and penetration-enhancing ability.

The selection of PTDs was based on their classification as primary, secondary, and nonamphipathic compounds. As a primary amphipathic PTD, transportan was selected. This type of PTD generally contains 20 or more amino acids and has sequential cationic and hydrophobic residues and affinity for neutral and anionic lipid membranes. Transportan seems to bind and insert into membranes, reduce surface tension at certain concentrations, and even induce phase transformation. Penetratin, representative of secondary amphipathic PTDs, was used due to its penetration-enhancing effect demonstrated by our group and others. Secondary amphipathic PTDs display their amphipathic properties after interacting with certain lipids and adoption of a helical structure separating charged and uncharged residues. TAT, one of the most studied PTDs, was selected to represent nonamphipathic peptides. These compounds generally have a large number of cationic amino acid residues and bind to negatively charged sulfated glycosaminoglycans of the surface of mammalian cells at low micromolar concentrations, followed by clustering and endocytic uptake. Macropinocytosis has been demonstrated to mediate TAT cell entrance.

Our results demonstrate that the nature of PTD plays an important role in the penetration-enhancing properties of topical nanocarriers. Among the PTDs studied, addition of transportan optimized drug cutaneous localization without compromising formulation safety. The transportan-containing ME improved drug efficacy against cutaneous tumor cells and three-dimensional (3D)-bioengineered skin cancer models, indicating the potential clinical usefulness of the nanocarrier for the treatment of cutaneous tumors.

Material and methods

Materials

BRJ 97 (polyoxyethylene 10 oleyl ether) and propylene glycol were obtained from Sigma-Aldrich Co. (St Louis, MO, USA). Monocaprylin was kindly supplied by Abitec Corporation (Janesville, WI, USA), and myvacet oil (diacetylated monoglycerides from soybean oil) was obtained from Quest (Norwich, NY, USA). Acetonitrile, ethanol, and methanol were purchased from Mallinckrodt Baker (Phillipsburg, NJ, USA), and paclitaxel from Polymed Therapeutics (Houston, TX, USA). Penetratin, transportan, and TAT were purchased from Anaspec (Fremont, CA, USA).

Methods

Preparation of MEs

The ME used in this study was previously developed by our group based on the water titration method. A surfactant mixture was composed of BRJ97:ethanol:propylene glycol at 2:1:1 (w/w/w) and was mixed with monocaprylin (oil phase) at 1.3:1 ratio, followed by addition of the aqueous phase at 43%. This concentration of water has been demonstrated to be sufficient to create an oil-in-water system. The pH of the formulations was kept between 5 and 5.5, because the skin surface pH ranges between 5 and 5.9, and formulations displaying pH closer to this range are generally considered safe.

Paclitaxel was incorporated in the oil phase to obtain a final concentration of 0.5% (w/w), whereas penetratin, transportan, or TAT was added to the aqueous phase to obtain a final concentration of 1 mM. Other peptides were able to increase drug penetration when not covalently attached at this concentration, which justifies its selection. It is important to note that, in this study, PTDs were mixed within the ME; it was envisioned that PTDs would function as penetration enhancers and diffuse out of the formulation and into the skin to modulate the barrier function. This approach differs from using noncovalent interactions to modify the surface of nanocarriers with PTDs, such as using metal chelating lipids that take advantage of interactions between chelated divalent metal ions and a short sequence of histidine residues added to the N- or C-terminus of proteins. Interaction of PTDs with the oil–water interface of ME droplets might be possible depending on peptide concentration and amphiphilic properties. Penetratin, for example, seemed to intercalate between monoolein hydroxyl headgroups and promote shrinkage in the reversed hexagonal phase lattice parameter.

The MEs containing paclitaxel and either TAT (METAT), penetratin (ME-P), or transportan (ME-T) were characterized for their isotropy using polarized light microscopy.
microscopy (Axiotop; Zeiss, Thornwood, NY, USA). Internal phase diameter and zeta potential were determined using a Zetasizer nanoseries instrument (Malvern Instruments, Malvern, UK) at room temperature.

In vitro skin penetration
Skin penetration of paclitaxel from the selected MEs was studied using Franz diffusion cells (diffusion area of 1 cm²; Laboratory Glass Apparatus, Inc, Berkeley, CA, USA), with porcine ear skin as model tissue and a receptor phase (3 mL) that consisted of 100 mM phosphate buffer (pH 7.4) containing 20% of ethanol (v/v) maintained at 37°C under constant stirring. Ethanol was added to aid drug solubility in aqueous medium as previously described.8 MEs (100 mg) were placed in the donor compartment of diffusion cells for 1–12 hours. At the end of the experiment, skin samples were rinsed with water to remove excess formulation, and the SC was separated from the epidermis (E) and dermis (D) by tape stripping. Fifteen pieces of tape were used; the first was discharged and the others were placed in conical tubes containing 4 mL of methanol for drug extraction.8 The remaining skin (viable epidermis + dermis, ED) was cut in small pieces, placed into conical tubes containing 2 mL of methanol, homogenized using a tissue homogenizer (Biospec Products, Bartlesville, OK, USA), and bath sonicated for 20 minutes. Using this method, drug recovery from skin layers was within 85%-93%.7 Paclitaxel delivered into SC, ED, and receptor phase was assayed by high-performance liquid chromatography (HPLC) after sample filtration (0.45 µm pore PTFE membranes). The assay was performed using a Shimadzu Prominance HPLC system equipped with a pump model LC-20AB, an autosampler model SIL-20A, a photodiode array detector model SPD-M20A set at 228 nm, and a Phenomenex C18 column (maintained at 25°C). The mobile phase was composed of 6:4 (v/v) acetonitrile:water at a flow rate of 1.0 mL/min.

To investigate whether delivery of paclitaxel occurred throughout the surface of the skin or was limited to certain structures the distribution of a fluorescent derivative of paclitaxel (Flutax, an Oregon Green 488 conjugate; Thermo Fisher Scientific, Waltham, MA, USA) in the skin was studied using fluorescence microscopy. The fluorescent derivative was incorporated into ME-T (since it delivered the largest amount of paclitaxel, as detailed in the “Results” section). After a 12-hour application of 100 mg of the fluorescent drug-loaded formulation, the diffusion area of the skin was carefully cleaned, frozen, embedded in Tissue-Tek OCT compound (Pelco International, Redding, CA, USA) and sectioned at 14 µm. The sections were analyzed under a 20× objective in a fluorescence microscope equipped with a filter for fluorescein isothiocyanate (Olympus Corporation, Tokyo, Japan). Time of exposure was fixed to minimize interference of autofluorescence, which was assessed with untreated skin sections.

ME effect on the skin barrier
Transdermal water loss was assessed as an index of the ability of MEs to induce changes in the cutaneous barrier.7,28 Transdermal water loss was measured before and after treatment with unloaded (not containing drug) plain ME, ME-P, ME-T, ME-TAT, or water using a closed chamber evaporimeter (Vapometer; Delfin Technologies Ltd., Kuopio, Finland) equipped with an adaptor to fit the diffusion cell (diffusion area of 1 cm²) opening.28 Skin sections mounted in diffusion cells were treated with 150 mg of the formulations for 5 minutes or 8 hours. The 8-hour period was selected as it was the earliest time point at which differences in the barrier-disrupting effect of solutions and MEs could be observed.28 After each period, skin samples were carefully wiped with tissue paper to remove the formulations, replaced back on the diffusion cells, and left undisturbed for 15 minutes before measuring the transdermal water loss (10 seconds).30 The skin was not rinsed since previous results demonstrated the influence of this procedure on transdermal water loss.31,32 The measurements at 5 minutes were performed to account for any effect of tissue wiping on the integrity of the skin barrier, and results were expressed as Δ transdermal water loss, calculated as the difference in transdermal water loss values after treatment for 8 hours and 5 minutes.

Influence of PTD on the irritation potential of formulations
In an attempt to differentiate the potential irritation induced by ME-T, ME-P, and ME-TAT, reconstructed human epidermis (MatTek Corporation, Ashland, MA, USA) was subjected to treatment with these formulations in comparison to PBS (negative control) or Triton (positive control) (25 mg). The protocols using human cells were considered as exempt for requiring approval by the Institutional Review Board of Albany College of Pharmacy and Health Sciences since sources are publicly available. Tissues were incubated at 37°C and 5% CO₂ for 2 hours, 5 hours, or 12 hours. After each treatment period, tissues were rinsed with PBS and incubated with 300 µL of 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide (MTT) solution (1 mg/mL) for 3 hours at 37°C and 5% CO₂ to account for tissue viability. MTT was then extracted by immersing the tissues in 2 mL of extracting solution (provided by the MatTek Corporation and a Phenomenex C18 liquid chromatography (HPLC) after sample filtration (0.45 µm pore PTFE membranes). The assay was performed using a Shimadzu Prominance HPLC system equipped with a pump model LC-20AB, an autosampler model SIL-20A, a photodiode array detector model SPD-M20A set at 228 nm, and a Phenomenex C18 column (maintained at 25°C). The mobile phase was composed of 6:4 (v/v) acetonitrile:water at a flow rate of 1.0 mL/min.

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with the extracted samples was determined at 570 nm (background reading at 650 nm was subtracted from the readings). Tissue viability (%) was plotted as a function of time.

To further differentiate the irritation potential among the formulations, extracellular interleukin-1α (IL-1α) levels were assessed in the culture medium using enzyme-linked immunosorbent assay (ELISA) (Thermo Fisher Scientific) according to the manufacturer’s instructions. This cytokine was selected because its rise is consistent with induction of an inflammatory cascade by irritants.31

Cytotoxic effect of ME-T containing paclitaxel against basal cell carcinoma culture

Based on its superior ability to promote paclitaxel localization in the skin without compromising safety, ME-T was selected for subsequent studies aimed at assessing formulation efficacy.

In this experiment, the cytotoxicity of ME-T against basal cell carcinoma cells was evaluated as an index of efficacy. Basal cell carcinoma cells were obtained from American Type Culture Collection (ATCC, Manassas, VA, USA) and grown at 37°C and 5% CO₂ atmosphere in Dulbecco’s Modified Eagle’s Medium (ATCC) containing 10% fetal bovine serum (Thermo Fisher Scientific) and additional penicillin and streptomycin (1%). For the cellular viability assay, cells were plated in 96-well plates (6,000 cells/well) and treated with either ME-T or propylene glycol containing paclitaxel diluted in cell culture medium (10–100 µg/mL) for 24 hours. Consideration that paclitaxel is a cycle-specific drug (cells are arrested in G2/M phase), an extra step was performed before the viability assessment to ensure time for the drug to act and to maximize cytotoxicity: following treatment, cells were rinsed with PBS and incubated in fresh medium (without treatment) for 36 hours. In previous studies, cells were maintained for periods of up to 6 days to maximize cytotoxicity. Cell viability after treatment with unloaded propylene glycol and ME-T diluted in cell culture medium (10–100 µg/mL) was also assessed to account for the effects of unloaded formulation.

Viability was determined using a cell proliferation assay reagent (CellTiter 96 Aqueous One Solution; Promega Corporation, Fitchburg, WI, USA). Cell culture medium (100 µL) containing 20 µL of the cell proliferation assay reagent was added to each well, and after incubation for 2 hours at 37°C and 5% CO₂, absorbance was recorded at 490 nm using a plate reader (SpectraMax; Molecular Devices LLC, Sunnyvale, CA, USA).

Cytotoxic effect of ME-T containing paclitaxel against reconstructed skin models of melanoma

To verify whether ME-T would be a viable strategy for treatment of skin tumors, its efficacy against more complex 3D models of the disease was assessed. The only commercially available bioengineered model is a melanoma model. It is acknowledged that the melanoma model presents distinct characteristics compared to nonmelanoma tumors (including basal cell carcinoma, squamous cell carcinoma) and other pretumor lesions that are considered treatable by topical formulations (such as low risk, superficial disease, and when the patient is a poor surgical candidate). However, use of this model was valuable as it allowed us to evaluate whether paclitaxel penetrates into the tissue at sufficient amounts to cause cytotoxic effects. Reconstructed skin models of melanoma (MLNM-FT-A375; MatTek Corporation) were incubated at 37°C and 5% CO₂ and treated with ME-T and myvacet oil (control vehicle) either unloaded or loaded with paclitaxel (25 mg) at 0.5% (w/w) every 3 days for 2 hours during a total period of 12 days. In this regimen, after the 2-hour treatment period, tissues were rinsed with PBS and incubated without formulation until the next treatment. The 2-hour treatment period was chosen as unloaded ME-T induced no significant reductions in tissue viability compared to PBS (see Results, item “Irritation potential of PTD-containing formulations”). The viability of the tissues was determined using MTT as described in the “Influence of PTD on the irritation potential of formulations” section.

In an attempt to establish a relationship between tissue concentration of the drug and effect, paclitaxel was quantified in a separate set of tissues submitted to the same treatment scheme. At the end of the experimental period, tissues were rinsed with PBS, homogenized with 1.5 mL of methanol, and the drug was assayed by HPLC as described in the “In vitro skin penetration” section.

Statistical analyses

The results are reported as mean ± standard deviation (SD). Data were statistically analyzed using the analysis of variance test followed by Tukey’s post hoc test (GraphPad Prism software). Values were considered significantly different when P < 0.05.

Results

Formulation characterization

PTD addition did not preclude ME formation, and all systems were fluid and isotropic as observed by polarized light microscopy. Droplet size was in the nanometer range, varying between 32.3 nm and 40.7 nm (Table 1, polydispersity index <0.27), and all formulations displayed a
Table 1 Characteristics of the microemulsions

<table>
<thead>
<tr>
<th>Formulation</th>
<th>Size (nm)</th>
<th>Zeta potential (mV)</th>
</tr>
</thead>
<tbody>
<tr>
<td>ME-T</td>
<td>32.3±2.9</td>
<td>4.1±0.7</td>
</tr>
<tr>
<td>ME-P</td>
<td>32.7±2.8</td>
<td>4.9±0.2</td>
</tr>
<tr>
<td>ME-TaT</td>
<td>40.7±6.5</td>
<td>6.8±1.2</td>
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Note: Values are expressed as average ± SD of three measurements.

Abbreviations: ME-T, microemulsion containing transportan; ME-P, microemulsion containing penetratin; ME-TaT, microemulsion containing TAT; SD, standard deviation.

small positive charge, with TAT-containing formulations displaying the highest value of zeta potential (+6.8 mV).

As previously reported, the plain ME displayed a similar size (39.5±2.1 nm) but slight negative zeta potential (−1.8±0.1 mV), demonstrating that PTD addition imparted a small positive charge in the system. All PTD-containing formulations were cationic, and thus, differences among the nature of charge (cationic versus anionic) should not interfere with formulation-mediated delivery as suggested by Baspinar and Borchert.

In vitro skin penetration

Independent of the presence and type of PTD, paclitaxel was detected mostly in the SC at 1 hour, suggesting a fast penetration in this layer. Skin penetration increased with time, and significantly larger amounts (P<0.01) of drug were quantified in the SC and viable layers of the skin (ED) at 12 hours compared to 1 hour for all formulations (Figure 1).

Comparing the penetration mediated by PTD-containing MEs with the plain formulation, only ME-T increased paclitaxel delivery into ED at both 6 hours and 12 hours in a significant manner (P<0.05). At 12 hours, ME-P also promoted a significant (P<0.05) increase in paclitaxel delivery to ED, while ME-TaT failed to do so. Among the PTD-containing formulations, paclitaxel skin (SC + ED) penetration at the longest time point studied (12 hours) was maximized with ME-T, whereas ME-TaT provided the lowest penetration (1.6-fold less). ME-T-mediated paclitaxel penetration into ED was significantly (P<0.05) higher at both 6 hours and 12 hours (1.6- to 1.8-fold) compared to ME-TaT. ME-P-mediated delivery was intermediate, and no significant difference (P>0.05) was observed compared to the either ME-T or ME-TaT at any of the time points studied.

Taken together, these results support previous observations that, although the presence of positive charge in nanocarriers has been associated with improved penetration, it may not be the only factor influencing the penetration-enhancing effect of the studied MEs. The nature of the PTD seems to play a role. When it comes to transdermal delivery, there was no significant difference among the formulations (delivery of paclitaxel varied between 0.36 µg/cm² and 0.44 µg/cm²).

To better understand the influence of the various PTDs on the kinetics of drug transport into the skin, we evaluated the rate of paclitaxel penetration into the SC and viable skin...
layers (Figure 2). Considering penetration into SC, all formulations displayed a maximum rate at 1 hour, suggesting fast drug penetration into this layer. ME-P and ME-T displayed similar maximum rates, while lower values were displayed by ME-TAT and the plain ME. Maximum rates of paclitaxel penetration into viable skin layers were achieved at a longer period of time (6 hours), with ME-T displaying the highest rate (1.75 µg/cm²/h). These results suggest that the presence and nature of the PTD do not influence the time necessary to achieve the maximum rate of paclitaxel penetration; however, they influence the maximum rate value obtained, especially into viable layers. Transportan-containing MEs displayed the most pronounced effect.

Considering that ME-T delivered the largest amounts of paclitaxel, drug distribution in the skin promoted by this formulation was studied using fluorescence microscopy. After administration of ME-T containing the fluorescent derivative of paclitaxel for 12 hours, the fluorescence seemed fairly homogenously dispersed on the tissue surface and viable skin layers, suggesting that ME-mediated delivery is not limited to certain skin structures/shunts but may occur throughout the SC (Figure 3).

**ME effect on the skin barrier**

To investigate whether the superiority of transportan-containing ME to improve paclitaxel penetration could be attributed to a stronger disruption in the skin barrier, the transepidermal water loss was evaluated and compared with water and the other formulations (Figure 4). Compared to water, all MEs increased transepidermal water loss, but the effect of ME-T was the most pronounced ($P<0.001$). These results suggest that transportan addition increased the disruptive effect of the ME more strongly. Additionally, the results also suggest that formulation-induced alterations in the barrier function could be detected at an earlier time point compared to a previous

**Figure 2** Rate of paclitaxel penetration into the skin.

**Note:** Data presented as average ± SD of three to seven replicates.

**Abbreviations:** SC, stratum corneum; h, hours; ED, viable epidermis and dermis; ME, microemulsion; ME-P, microemulsion containing penetratin; ME-T, microemulsion containing transportan; ME-TAT, microemulsion containing TAT; SD, standard deviation.

**Figure 3** Fluorescence microscopy of skin sections treated with ME-T containing the fluorescent derivative of paclitaxel.

**Note:** The images show untreated skin observed under fluorescent light to show interfering fluorescence (A) and skin treated with paclitaxel in ME-T (B).

**Abbreviation:** ME-T, microemulsion containing transportan.

**Figure 4** Difference in transepidermal water loss after treatment of skin sections (Δ transepidermal water loss) for 5 minutes or 8 hours with water (control) and microemulsions containing PTDs.

**Notes:** Data presented as average ± SD of six to seven replicates. *$P<0.05$ compared to water, **$P<0.01$ compared to water, and ***$P<0.001$ compared to water.

**Abbreviations:** ME, microemulsion; ME-T, microemulsion containing transportan; ME-P, microemulsion containing penetratin; ME-TAT, microemulsion containing TAT; PTDs, protein transduction domains; SD, standard deviation.
Irritation potential of PTD-containing formulations

We next evaluated whether the type of PTD influenced the irritation potential of the MEs. PBS is considered safe and, as previously observed, did not reduce tissue viability during the time period studied (Figure 5A). Compared to PBS, the viability of the tissues treated with Triton was significantly ($P<0.05$) reduced to 73.9%±4.5% after 2 hours, while a longer period of time (5 hours) was required for the PTD-containing MEs to display similar effects. At 5 hours, Triton-treated tissues displayed significantly lower ($P<0.05$) viability than ME-treated tissues, suggesting the lower irritation potential of the formulations. Comparing the MEs among each other, ME-T-treated tissues displayed lower, but not significantly different, viability at 12 hours compared to ME-P or ME-TAT.

IL-1$\alpha$ is believed to be one of the cytokines involved in the initiation of the inflammatory cascade in response to skin exposure to irritants. Measurements of IL-1$\alpha$ released from keratinocytes of epidermal skin equivalents have been used to predict and rank the irritation potential of formulations. We assessed the extracellular levels of this cytokine in an attempt to further differentiate the influence of the various PTDs on the formulation irritation potential (Figure 5B). At the longest time point studied (12 hours), ME-T promoted the most pronounced increase in IL-1$\alpha$ release among the MEs (1.8- and 1.5-fold compared to ME-P and ME-TAT, $P<0.05$), but these cytokine levels were still 5.8-fold lower than those induced by Triton. These results suggest that the transportan-containing ME displays the most pronounced irritation potential among the MEs, which is not unexpected since it also presented a stronger ability to increase transdermal water loss. However, given the fact that its effects on IL-1$\alpha$ release were less than those of Triton, it is reasonable to suggest that ME-T should be better tolerated by the skin, and thus, addition of transportan does not compromise formulation safety.

Cytotoxicity of ME-T containing paclitaxel against basal cell carcinoma culture

Since ME-T maximized the cutaneous delivery of paclitaxel and was safer than the moderate irritant Triton, the efficacy of this formulation against basal cell carcinoma cells was studied and compared to a drug solution in propylene glycol. Cell viability after treatment with the unloaded formulations at any of the concentrations used was at least 80% (Figure 6). Paclitaxel-loaded solution and ME-T led to significant reductions in cell viability ($P<0.05$) compared to the unloaded formulations, suggesting that the stronger cytotoxicity of the loaded formulations derive from the presence of the drug and that incorporation in the ME did not hinder drug activity.

Drug cytotoxicity was dose dependent. When the concentration of paclitaxel solution in the treatment medium was 10 $\mu$g/mL (and drug concentration was 0.05 $\mu$g/mL, cell viability was 80.4%, which represents a reduction of 16% compared to the unloaded vehicle. Increasing the concentration of drug solution to 100 $\mu$g/mL (and drug...
concentration to 0.5 µg/mL) further reduced cell viability to 59.4% (P<0.05). Compared to the drug solution, treatment with paclitaxel-loaded ME-T promoted a curve shifting, ie, treatment with the same concentration of the drug-loaded ME resulted in lower viability. At the largest concentration (100 µg/mL), ME-T promoted a twofold reduction in cell viability (P<0.05). Considering that there was no significant difference in cell viability when comparing the unloaded formulations at the concentrations used, the more pronounced reduction in cell viability provided by the loaded ME-T may potentially result from a more efficient drug delivery and higher uptake of the drug.

Cytotoxicity of ME-T containing paclitaxel against reconstructed skin models of melanoma

Having demonstrated the cytotoxic effects of loaded ME-T against tumor cells, we next evaluated the formulation efficacy against a reconstructed skin model of melanoma (MLNM-FT-A375; MatTek Corporation). Similar to what we observed in cells, tissue viability after treatment with paclitaxel-containing ME-T was significantly lower (2.2-fold, P<0.005) than that with the unloaded ME (Figure 7), demonstrating that the presence of paclitaxel promoted a more pronounced decrease in tissue viability.

To verify whether drug incorporation into ME-T increased cytotoxicity compared to simple solutions, tissue viability after treatment with paclitaxel-loaded myvacet oil solution at the same concentration was assessed. As can be observed in Figure 7, cell viability was approximately twofold lower after treatment with ME-T (67.4±5.8% and 34.5±2.9% for myvacet solution and ME-T, respectively), demonstrating the benefit of the nanocarrier.

To establish a relationship between reduction in cell viability and drug delivery to the skin, paclitaxel concentration in the tissue was assessed. Paclitaxel delivery into the skin after treatment with ME-T was 4.5-fold higher (P<0.005) compared to the solution. These results demonstrated that the more pronounced reduction in tissue viability mediated by ME-T resulted from a larger amount of drug delivered into the skin.

Discussion

The interest in topical formulations aimed at increasing localization of paclitaxel in the skin has grown over the past years. In a previous study, we assessed the skin penetration of paclitaxel from cationic MEs containing either the PTD penetration or the lipid phytosphingosine. In that study, only one PTD (penetratin) and one time point (12 hours) were evaluated. The superiority of the penetratin-containing ME encouraged us to compare the influence of PTDs from various classes on the skin permeability, penetration kinetics, and formulation irritation as described in this study. Furthermore, in this study, we advanced in formulation evaluation and assessed the potential clinical usefulness of a selected system by studying its efficacy against cutaneous tumor cells and a bioengineered 3D melanoma model.

Compared to other formulations studied, MEs have the advantage of easy preparation, thermodynamic stability,
and versatility of composition, which can reduce costs. However, due to the high concentration of surfactants and addition of penetration enhancers (including the PTDs), MEs may display increased irritation potential. Thus, one of the goals of this study was to verify whether MEs containing PTDs would be tolerated by the skin. The toxicity of PTDs has been well studied in various cell lines, but to the best of our knowledge, this is the first report comparing the effects of PTDs from different classes on formulation-induced irritation in bioengineered skin. Among the formulations, the most pronounced effect on IL-1α release was displayed by ME-T. Since this effect was less pronounced than that induced by the moderate irritant Triton, it is reasonable to suggest that the formulation should be better tolerated by the skin, and thus, transportan addition does not compromise formulation safety.

The increase in the IL-1α release mediated by transportan compared to the other PTDs was counterbalanced by its ability to maximize paclitaxel cutaneous delivery. These effects may be related to the more pronounced ability of the formulation to disrupt biological barriers, as demonstrated by the increased transepidermal water loss among all studied MEs. The TAT-containing ME displayed the least effect. The following two conclusions can be drawn from these results: first, as previously observed, the presence of positive charge in nanocarriers does not seem to be the only relevant factor affecting paclitaxel delivery as all PTD-containing MEs displayed similar slight positive charges. Second, not all PTDs might penetrate the skin and function equally like penetration enhancers in a given condition. Considered a primary amphipathic PTD, transportan is long enough to theoretically span the hydrophobic core of bilayers. Occurrence of a connection between the two monolayers through the peptide might lead to bilayer thinning and formation of pores. These effects provide reasonable justification for the enhanced ability of ME-T to decrease skin barrier function, while increasing IL-1α release and paclitaxel penetration. Previous studies have suggested a correlation between amphiphilic properties and toxicity of PTDs.

ME-T improved paclitaxel cytotoxicity against cultured basal carcinoma cells compared to a control drug solution, most likely due to a more efficient delivery. This result is consistent with previous reports demonstrating the efficacy of nanocarriers to improve drug cytotoxicity against cutaneous tumor cells. The stronger cytotoxic effect of paclitaxel-containing ME-T was also observed in melanoma tissues, which displayed viability <40%. This low viability suggests that paclitaxel affects not only cancer cells but also normal cells in the tissue, which is in agreement with other studies demonstrating the cytotoxic and proliferation-inhibiting effects of paclitaxel against several cell types, including normal cells. This wide-range cytotoxicity reinforces the importance of localizing paclitaxel in the lesions to prevent adverse effects in other tissues/organs. This effect was associated with a higher drug concentration in the tissue, demonstrating the relationship between increased cytotoxicity and enhanced drug delivery by ME-T. It is worth pointing out that formulation-mediated penetration would be more pronounced in the melanoma tissue than in porcine ear skin (the sum of drug in SC and viable epidermis) if we take into consideration the time of application, which is consistent with previous studies demonstrating a higher permeability of bioengineered tissues compared to the porcine and human skin. Higher permeation flux values of 500–800 times than those for human skin have been reported for skin equivalent models compared to human skin, and this difference seems to be more pronounced for lipophilic compounds.

PTDs have been most frequently studied for systemic delivery, but the high in vitro uptake rates, the relatively low specificity, and the fast clearance rates observed in several studies suggest that their topical application presents a higher potential. Our results support the potential of selected PTDs to improve drug topical delivery, leading to a more pronounced efficacy against a bioengineered model of melanoma. Given the fact that the PTDs were simply mixed into ME, their amphiphilic properties may play an important role as demonstrated by the stronger effect of ME-T compared to ME-TAT, at least within the experimental conditions used here. Although we emphasized the effect of the PTD on the penetration-enhancing ability of the ME, it is necessary to consider the ME role on the PTD penetration. As demonstrated by Kim et al, the pore-forming peptide magainin alone was not able to affect skin permeability, likely due to its relatively large size and insufficient penetration into the tissue. Skin permeability to fluorescein was increased by 47-fold when magainin was combined with N-lauroyl sarcosine in a 50% ethanol solution, which indicates that the use of formulations capable of promoting peptide penetration aids its effect on skin permeability.

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**Disclosure**

The authors report no conflicts of interest in this work.

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