Original Research Article

Potentiation of raloxifene cytotoxicity against MCF-7 breast cancer cell lines via transdermal delivery and loading on self-emulsifying nanoemulsions

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Abstract

**Purpose:** To enhance raloxifene (RLX) delivery and cytotoxicity against breast cancer (MCF-7) cell lines.

**Methods:** This was a solubility study of RLX in different oils, surfactants, and co-surfactants. Twelve formulae were tested to reach the smallest globular size, and hydroxypropyl methylcellulose (HPMC), and Carbopol 947 polymers were tested for formation of transdermal films. The formula with the lowest size was compared with raw RLX in diffusion studies using a Franz diffusion cell. Finally, a cytotoxicity study against MCF-7 breast cancer cell lines was conducted.

**Results:** The maximum solubility of RLX was in Tween 80, peppermint oil, and PEG 200; therefore, these were the main components of the 12 formulations. The release of RLX loaded on the self-nanoemulsion drug delivery system (SNEDDS) was increased 3-fold compared with raw RLX. Cytotoxicity results revealed that RLX SNEDDs decreased MCF-7 cell survival by approximately 40%, compared with raw RLX (control), which augmented the RLX suppression of breast cancer cell lines.

**Conclusion:** Improvement in RLX cytotoxicity is a novel strategy to suppress breast cancer.

**Keywords:** Raloxifene, Osteoporosis, Bioavailability, Nanoemulsion, Nanoparticles

INTRODUCTION

Raloxifene (RLX) is used to treat osteoporotic postmenopausal women and may also lower the chances of acquiring certain types of breast cancers (i.e., invasive breast cancer) [1]. RLX is not an oestrogen hormone, but it acts like oestrogen in some parts of the body. During menopause, oestrogen levels in the blood decrease, affecting bone cells and tissues. RLX acts like an oestrogen blocker [2]. Further, RLX does not relieve menopausal symptoms such as hot flashes, as it belongs to a class of drugs known as selective oestrogen receptor modulators (SERMs). The main problem is to improve its low bioavailability (2%) resulting from excessive hepatic metabolism [3].

Nanoemulsions consist of nanosized droplets with a diameter of < 300 nm, typically 20–200 nm. The thermodynamically non-balance of these transparent, heterogeneous dispersions of
two non-mixtures fluids are found in [4]. Thanks to the long, almost several months, kinetic destabilization, they are kinetically stable. Nearly four decades ago, their bioavailability was investigated. Nanoemulsions are also referred to as thermodynamic solidity due to the presence of fine beads, which resist sedimentation, flocculation, and mixing due to Brownian motion [5,6]. These nanovehicles provide the benefits of high interfacial zones, straightforwardness of preparation, low thickness, and long-lasting colloidal states, both for hydrophilic and hydrophobic medications. Thus, they provide increased dependability, more prominent transmucosal and transdermal medication conveyance, and, consequently, enhanced bioavailability [7,8].

In addition, nanoemulsions convey noteworthy wetting, spreading, and entrance capacities and can be scaled up because of their simplicity of fabrication. Nanoemulsions are also strong, accounting for their effectiveness, which additionally enhances their usefulness. The bioavailability of the emulsions are enhanced incrementally when the bead sizes approach 100, 90, and 30 nm. The purpose of this study was to improve RLX bioavailability through the use of self-nanoemulsion drug delivery systems (SNEDDS) stacked in a transdermal location to maintain a strategic distance from digestive processes [9].

**EXPERIMENTAL**

**Materials**

RLX powder was a gift from EIPICO Pharmaceuticals (Cairo, Egypt); olive oil, castor oil, Tween 20 polysorbates, polyethylene glycol 200, glycerin, dill oil, ethyl alcohol, Tween 80, and sesame oil were purchased from Sigma-Aldrich (Dusseldorf, Germany).

**Solubility study**

The solubilities of RLX in various oils (olive, sesame, dill, castor, and peppermint), surfactants (Tween 80 and Span 80), and co-surfactants (PEG 200, ethyl alcohol, and glycerine) were determined by the addition of excess amounts of the drug to screw-capped glass vials containing 1 mL of each of the solutions. The mixtures were placed in a thermostatically controlled shaking water bath at 37 °C for 24 h. The vial contents were then examined for clarity; if it was clear, more drug was added. Then, the mixtures were centrifuged at 15,000 rpm for 1 h. The supernatant was diluted with methanol, and the concentration was determined by ultraviolet (UV) spectrophotometry at a wavelength of 289 nm.

**Preparation of nanoemulsion**

Based on the above-mentioned solubility studies, the selected oil (peppermint oil), surfactant (Tween 80), and co-surfactant (PEG 200) were chosen for preparation of nanoemulsions. Twelve formulae were prepared using varying concentrations of oil, surfactant, and co-surfactant with 30 mg of the drug, as shown in Table 1. For any mixture, the total weight of the three components was always maintained at 2 gm.

**Table 1: Composition of nano emulsion formulations**

<table>
<thead>
<tr>
<th>Batch</th>
<th>Peppermint oil (mg)</th>
<th>Tween 80 (mg)</th>
<th>PEG 200 (mg)</th>
<th>Globular size (nm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>F1</td>
<td>0.35</td>
<td>0.5</td>
<td>0.15</td>
<td>323</td>
</tr>
<tr>
<td>F2</td>
<td>0.35</td>
<td>0.55</td>
<td>0.1</td>
<td>234</td>
</tr>
<tr>
<td>F3</td>
<td>0.3</td>
<td>0.475</td>
<td>0.22</td>
<td>321</td>
</tr>
<tr>
<td>F4</td>
<td>0.3</td>
<td>0.55</td>
<td>0.15</td>
<td>543</td>
</tr>
<tr>
<td>F5</td>
<td>0.271</td>
<td>0.501</td>
<td>0.2</td>
<td>298</td>
</tr>
<tr>
<td>F6</td>
<td>0.25</td>
<td>0.55</td>
<td>0.2</td>
<td>651</td>
</tr>
<tr>
<td>F7</td>
<td>0.25</td>
<td>0.475</td>
<td>0.275</td>
<td>942</td>
</tr>
<tr>
<td>F8</td>
<td>0.2</td>
<td>0.55</td>
<td>0.25</td>
<td>267</td>
</tr>
<tr>
<td>F9</td>
<td>0.4</td>
<td>0.425</td>
<td>0.375</td>
<td>763</td>
</tr>
<tr>
<td>F10</td>
<td>0.15</td>
<td>0.475</td>
<td>0.375</td>
<td>176</td>
</tr>
<tr>
<td>F11</td>
<td>0.15</td>
<td>0.375</td>
<td>0.475</td>
<td>321</td>
</tr>
<tr>
<td>F12</td>
<td>0.1</td>
<td>0.35</td>
<td>0.55</td>
<td>421</td>
</tr>
</tbody>
</table>

**Globular size analysis**

A specific volume of each formula (100 µL) was poured into 50 mL of distilled water to allow SNEDDS to form emulsions and then directly subjected to a Microcrack Nanosizer (Microtrac, Montgomeryville, PA, USA) to measure globular size and zeta potential.

**Globular morphology**

The globular morphology of the selected formula was studied by transmission electron microscopy (TEM) to determine the globular size and morphology. TEM analysis (TEM Tecnai 200 Kv TEM Fei; Electron Optics, Tokyo, Japan) of the RLX SNEDDS was performed for morphological characterization and visualization of emulsion droplets. A drop of sample obtained after dilution was placed on copper grids, stained with 1% phosphotungstic acid for 30 s, and finally analyzed by TEM to visualize the droplet morphology.

**Transdermal patch formulation**

The selected formula of RLX SNEDDS was poured into a polymeric solution of 1% of HPMC,
HPC, and chitosan for the preparation of transdermal patches. All films used 1 g of each polymer in 100 mL tap water with 30 mg RLX powder as the control. Polymeric solutions were poured into petri dishes and covered with aluminum foil and then transferred to a 45°C oven until complete evaporation of the solvent was achieved. All films were characterized for folding properties, appearance, and film elongation, as shown in Figure 2 (RLX SNEDDS TEM morphology).

Preparation of RLX SNEDDs transdermal patch

The prepared polymeric solutions were stored in an oven at 45°C for almost three days. The four patches were not completely dry and had a bright yellow color. Before cutting the patches, the drug contents of a 1.5 cm two piece from each sample was calculated.

In vitro release studies

The prepared patches were tested for diffusion of drug using automated Franz diffusion cell apparatus (Microette plus, USA) with a diffusion area of 2.85 cm and 7 ml volume receptor chamber. Patches prepared with RLX, and the Formula 10 was used. In the receptor chamber, a buffer with the pH 7.2 was used as a medium at 32.5 °C ± 2.0, and the rate of stirring was 400 rpm. The auto sampler is programmed to take samples of the buffer solution at preselected times (0:30, 1, 2, 4, 6, 8, 12 h) with equal amounts to the receptor chamber. HPMC was chosen as a transdermal patch matrix.

Evaluation of RLX SNEDDs cytotoxicity against MCF-7 cells

MCF-7 breast cancer cells were gifted from Dr Serag El-bahiry’s, Cell Culture Lab, Faculty of Science, King Khalid University, Abha, KSA. The cells were grown in complete growth media composed of 5 % fetal bovine serum (FBS), 2 mM L-glutamine, 100 μg/mL streptomycin, 100 U/ml penicillin, and 2.2 g/l NaHCO3. Cells were maintained at 37 °C in a humidified atmosphere of 5% CO2. MCF-7 breast cancer cells were seeded in 96 well plates, incubated for 24 h at 37 °C in 5 % CO2, and then treated with RLX or RLX SNEDDs. DMSO (0.1 %) served as the vehicle control. The cells were incubated for 72 h and fixed using a trichloroacetic acid (TCA) 10% solution. Cell number was determined using the sulforhodamine B (SRB) assay. The concentration required to decrease cell number by 50 % (IC50) was determined by nonlinear regression using Prism software. For the time-course analysis, cells were treated with RLX or RLX SNEDDs (0.01 – 100 μg/mL). Three independent experiments were performed in triplicate.

RESULTS

We experimented with the oils to see which one has the highest solubility of RLX and found that the peppermint oil has the highest RLX solubility with a concentration of 1.69 mg/mL. The highest solubility was found with the surfactants and co-surfactants; further, Tween 80 had the highest of the surfactants with a concentration of 3.703 mg/mL, and PEG 200 had the highest of the co-surfactants with a concentration of 4.917 mg/mL, as shown in Figure 1.

In vitro drug release

With regard to RLX SNEDDS morphology, TEM images show circular globules with size less than 1 µm; this spherical globule facilitates skin penetration and homogenize rate of diffusion, as shown in Figure 2.
The prepared patches were tested for diffusion of the drug using an automated Franz diffusion cell apparatus (Microette™ Plus, USA) with a diffusion area of 2.85 cm² and a 7 mL volume receptor chamber. The patches prepared with RLX and Formula 10 were used. In the receptor chamber, a buffer (pH 7.2) was used as a medium at 32.5 ± 2.0 °C, and the rate of stirring was 400 rpm. The autosampler was programmed to take samples of the buffer solution at preselected times (0 and 30 min, and 1, 2, 4, 6, 8, and 12 h) with equal amounts to the receptor chamber. HPMC was chosen as a transdermal patch matrix.

Figure 3: % of RLX cumulative released after 12 h. (O) refer to RLX transdermal patch while (O) refers to Raw -RLX

To provide the potency of RLX SNEDDS raw RLX in the MCF-7 breast cancer cell lines, IC50 curves were initially produced. Both raw RLX and RLX SNEDDS demonstrated a consistent cytotoxicity across the cell lines, but RLX SNEDDS was more potent than raw RLX. Specifically, IC50 values for RLX SNEDDS was 2.77 ± 0.38 μM, while the raw RLX was 1.98 ± 0.17 μM, as shown in Figure 4.

Figure 5: IC50 (% of Raw RLX and RLX SNEDDs. (O) denote RLX SNEDDS while (O) = Raw -RLX

Nanoemulsions are structures that are thermodynamically stable and grow when a particular solution quantity is poured into liquid. When the peppermint oil concentration decreased from Tween 80 to PEG 200[4], a higher region of nanoemulsion was found, and about 35% of dill oil was solubilized. The interfacial tension may be further reduced by increasing the interface fluidity; therefore, the oil stage may further penetrate the hydrophobic area of 80 monomer. The inclusion of PEG 200 as a co-factant minimized the interface end tension and rendered the interface film versatile enough to show multiple curvatures needed to form nanoemulsion under a range of circumstances. The nanoemulsion region decreased when the amount of PEG200 was raised in contrast to Tween 80, leading to the creation of a smaller quantity of micelles [10].

The improved cell cytotoxicity and penetration of SNEDD RLX can be used to improve raw RLX dose effectiveness against cancer cells MCF-7, which could reduce RLX adverse events [11–13]. SNEDDs are taken by human cells through a number of concurrent pathways. SNEDDs increase RLX inclusion and enhance mobile RLX absorption by attracting a cell membrane negatively charged and activating a receptor-free cascade, contributing to the incorporation of RLX into the cellular cytoplasm. The interaction between SNEDDs and the cell membrane includes the lure of proteoglycans and surfactants. Therefore, SNEDDs activate intracellular signalling cascades, which enhance the organic cycle of absorption.

SNEDDs can penetrate cells using distinctive endocytic pathways such as pinocytosis along with macropinocytosis, clathrin-mediated endocytosis, caveolae or Q7 lipid-raft-mediated endocytosis, and clathrin-or caveolae-impartial endocytosis. Endocytic uptake pathways rely on peptide characteristics, load characteristics of SNEDD, and specific cell characteristics. Transport of RLX may be controlled with the aid of the extracellular carrier system’s metabolism, which involves peptide uptake, and intracellular degradation in endocytic vesicles. Endosomal escape could help to prevent degradation of RLX in lysosomes and thus aid the delivery of RLX to its biological target [14–16].

CONCLUSION

A new dosage form of RLX as a transdermal patch to overcome extensive hepatic first-pass metabolism of oral dosage using SNEDDs was established. After conducting in vitro release studies of films, the results showed that RLX
transdermal patches demonstrated a > 3-fold release of RLX compared with raw RLX, which suggested a significant enhancement in RLX bioavailability to improve cytotoxicity. This novel method could potentially be used for suppressing breast cancer cells.

DECLARATIONS

Conflict of interest

No conflict of interest is associated with this work.

Contribution of authors

We declare that this work was done by the authors named in this article and all liabilities pertaining to claims relating to the content of this article will be borne by the authors.

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