Formulation of Triple Cream for Treatment of Melasma

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Abstract

Background: Melasma is a common disorder of hyperpigmentation affecting the face that is associated with considerable psychological impacts. The management of melasma is challenging and requires a long-term treatment plan. Therapeutic goals for hyperpigmentation include the destruction of melanosomes, inhibition of the melanocyte formation, and delay in the spread of melanocytes. Hydroquinone is a commonly used treatment for melasma, but it is combined with other medications due to its low efficacy in single therapy and relapse. Hydroquinone is hydroxy phenol which inhibits melanin by inhibiting tyrosinase. In addition, retinoid is effective in treating melasma and stimulate the cycle of skin cells, thus increasing the rate of loss of color pigmentation. A corticosteroid is used to increase the efficacy of these two drugs.

Objectives: The purpose of this study is to formulate an integrated formulation for the treatment of this skin disease.

Methods: The triple cream, which is a combination of hydroquinone (HQ), tretinoin (TRE), and fluocinolone acetonide (FLU) demonstrate better efficacy in decreasing skin pigmentation due to the additive and synergistic effects of these ingredients. As there is no topical generic formulation in combined form in Iran, such cream is prepared that contains 4% HQ, 0.05% TRE and 0.01% FLU as an oil in water (O/W) cream.

Results: The result of this study was a 6-month accelerated evaluation of physical and physicochemical properties, including pH, electrical conductivity, density, viscosity, active ingredient identification, determination of amount and completely stable and homogeneous microbial tests. A review of the identification of the active ingredient and the determination of the amount conformed to the USP standard (American Pharmacopoeia). Also, material release from the base was well done.

Conclusions: The prepared cream was completely stable and homogeneous during the accelerated conditions by evaluating of physicochemical and microbial assessments.

Keywords: Fluocinolone, Hydroquinone, Melasma, Tretinoin

1. Background

Melasma is the overproduction of melanin in the skin and came from the Greek translate of phrase black (Melas). Melasma is one of the most popular, therapy-resistant types of acquired hyperpigmentation. The risk of melasma infliction in females is nine times more than males, and it is more often in darker-skinned individuals, such as Hispanics, Asians, or other racial groups (1). Melanocytes create melanin pigment and the increase in melanin may be due to both a proliferation in melanogenesis and melanocytosis. The contributing factors in melasma pathogenesis include genetics, hormones, and UV radiation. Other implicated factors include the use of certain cosmetics, phototoxic drugs, and anticonvulsant medication. Lots of long-term treatment plan of melasma shows non-effective procedure. Therefore, it is a challenging disease (2).

Bleaching creams are a common OTC or prescribed products used to whiten the skin color spots. These creams contain various active ingredients such as hydroquinone (HQ), glycolic acid, azelaic acid, kojic acid, corticosteroids, tretinoin (TRE) and lots of herbal extracts (3). Based on the cost/benefit analysis (4), the triple combination therapy is the most effective primary treatment for melasma rather than monotherapy (4). The combination of HQ, TRE, and topical steroids can provide better therapeutic outcomes (as they work with different mechanisms of inhibition of melanogenesis) and also can be tolerated by patients (5). The HQ is the main treatment for melasma but has relatively low effectively as monotherapy, and re-
currences are common. This ingredient was originally believed to act mainly by inhibition of tyrosinase of epidermal melanocytes because of its structural analogy to melanin precursors.

The whitening effect of HQ is seen after 2 to 3 weeks of daily use (6). Its efficacy as a depigmenting agent is influenced by several parameters, such as concentration, chemical stability, and vehicle of administration. The highest concentration of HQ is more effective, but it may cause dermatitis. Making HQ into a stable preparation is challenging because of the highly reactive oxidative nature of the agent, changing from creamy white to a darker brown via oxidation. The penetration of enhancers such as TRE or alpha hydroxyl acids enhances the timed delivery of HQ to the skin (7). The first time, TRE was used in hyperpigmentation by Kligman in 1975. His statement showed that patients who getting TRE developed lighter skin after months of applications. In fact, TRE acts by preventing the oxidation of HQ, improving stratum corneum penetration, allowing pigment elimination, and increasing keratinocyte proliferation. As a result, TRE is commonly combined with one or two other agents to accelerate the beneficial effect (8).

Triple combination also contains a corticosteroid. It is believed that corticosteroid can inhibit melanin synthesis by decreasing cellular metabolism. Furthermore, corticosteroids reduce the irritation or inflammation caused by HQ and TRE. In 1975, Kligman and Willis used 0.1% dexamethasone in combination with 5% HQ and 0.1% TRE (9). After that, scientists found out steroids containing fluorne were more effective than nonfluorinated ones. Therefore, 0.01% fluocinolone acetonide (FLU) was used instead of dexamethasone, then HQ and TRE were reduced to 2% - 4% and 0.05%, respectively (3). Now in the US, the pharmaceutical formulation of this triple-combination was sold under the brand name of Tri-Luma (Galderma Co.) (4, 10, 11). Although combination therapy shows an acceptable efficacy, multi-active formulas are unstable both in physical and chemical reactions. Parameters such as phase separation, solubility, oxidation, assay, content uniformity and etc. are important (12). To the best of our knowledge, there is no study about the formulation type and stability tests of triple bleaching creams. Neither method has been mentioned in the United States pharmacopeia. In this research, we formulated and examined physico-chemical properties of a triple combination of HQ 4%, TRE 0.05%, and FLU 0.01%.

Side effect of HQ is skin irritation, dryness, erythema, and dermatitis. Common side effects of topical TRE include warmth or stinging immediately after applying the medication. Other side effects of TRE include skin tingling, itching, redness, swelling, dryness, peeling, and irritation during the first 2 - 4 weeks of using the medication. Common side effects of FLU include burning, stinging, itching, dryness, peeling, and redness. These side effects should disappear in a few days as your body adjusts to FLU.

2. Methods

The HQ was obtained from Medilux (India), TRE was procured from Olon (Italy) and FLU was purchased from Farmabios (Italy). Glycerol stearate and PEG-100 stearate and caprylic/capric acid triglyceride were bought from Croda (UK), cetearyl alcohol, glycerin, methylparaben, propylparaben, sodium metabisulfite (MBS), and ascorbic acid were purchased from Merck (Germany). Deionized water was produced freshly whenever required. A lipid phase containing FLU, TRE, glycerol stearate and PEG-100 stearate, caprylic/capric acid triglyceride and cetearyl alcohol, totally 25%, as well as 75% aqueous phase composed of HQ, ascorbic acid, glycerin and methylparaben, propylparaben, MBS, and deionized water.

2.1. Preparation of Triple Cream

To prepare a triple cream, we used a lipid phase containing FLU, TRE, glycerol stearate and PEG-100 stearate, caprylic/capric acid triglyceride and cetearyl alcohol, as well as an aqueous phase composed of HQ, ascorbic acid, glycerin and methylparaben, propylparaben, and sodium metabisulfite (MBS), and deionized water. The Aqueous phase was heated up to 75°C ± 1°C. Oily phase was heated to the same temperature. Then oil in water (O/W) cream was prepared by the adding of the oily phase to the aqueous phase with nonstop stirring at 600 rpm (IKA, Germany) until it was cooled down to the room temperature. This formulation must be kept in the refrigerator at 5 ± 3°C (13, 14).

2.2. Stability Studies

The manufactured cream was passed 6 months accelerated stability tests at 25 ± 2°C/60 ± 5% RH. During this period, physical and chemical analysis was done at 1st, 2nd, 3rd, and 6th months as mentioned in ICH stability guideline (15).

2.3. Physical Parameters

The prepared cream was examined visually for color, odor, homogeneity, consistency, phase separation.

2.4. Emulsion Type

An electrical conductivity test was conducted by conductivity meter to determine the type of emulsion. In this test, the sample was diluted with deionized water. If the emulsion is completely soluble in deionized water and forms a diluted homogeneous emulsion, the emulsion is O/W and vice versa (16).
2.5. Density

The density, or more precisely, the volumetric mass density, of a substance is its mass per unit volume. The exact volume of pycnometer was determined by filling it fully with water and was tapped the sides gently to remove the air bubbles. The flask was weighted full of cream, then easily the density was calculated (dividing mass by volume).

2.6. Determination of pH

To this end, 2 ± 0.01 g of cream was dispersed in a 50-mL beaker containing 18 mL water. Then pH was determined in 25 ± 2°C by pH meter (Metrohm 827, Swiss) (17).

2.7. Viscosity

The viscosity of cream was measured by Polyscience Viscometer (Instrumentacion Cientifica Tecnica, Spain) using spindle R7 at 100 RPM with the run time of 1 minute. Viscosity determinations were performed at room temperature (25 ± 2°C) (17).

2.8. Identification of Active Ingredients

For identification of HQ, a portion of cream equivalent to about 50 mg of HQ was dissolved in a mixture of chloroform and methanol to make 50 mL dispersion, then was read in TLC paper and compared with the standard. The retention time of the major peak in the chromatogram of the assay preparation corresponding to that in the chromatogram of the standard preparation was obtained as directed in the assay of TRE. At least for FLU, a quantity of cream equivalent to about 0.5 mg of FLU was dissolved, and centrifuged with water and chloroform (1:2), then the aqueous layer was removed and sodium sulfate 100 mg/mL was added. The extract was responded to TLC and then compared with standard solutions.

2.9. Assay of Active Ingredients

2.9.1. Hydroquinone (HQ)

An accurately weighed portion of cream, equivalent to about 20 mg of HQ, was moved to a 100 mL beaker, which contained 50 mL methanol to make 50 mL dispersion, then was read in TLC paper and compared with the standard. Seventy-five mL of methanol was added to 25 mL of the sample solution, and centrifuged with water and chloroform (1:2), then the aqueous layer was removed and sodium sulfate 100 mg/mL was added. The extract was responded to TLC and then compared with standard solutions.

2.9.2. Tretinoin (TRE)

An accurately weighed amount of cream, equivalent to 1 mg of TRE, was moved to a 50 mL volumetric flask, and added 20 mL of tetrahydrofuran. 5 mL of this solution was transferred to a 25 mL volumetric flask, diluted with a mixture of tetrahydrofuran and diluting solution (prepared a mixture of water and dilute phosphoric acid 10%) (3:2) to volume. The separatists were injected equivalent volumes (about 25 µL) of the standard preparation and the assay preparation into the chromatograph, the chromatograms were verified at 352 nm.

2.9.3. Fluocinolone Acetonide (FLU)

An accurately weighed portion of cream, equivalent to about 0.75 mg of the FLU, was dissolved in about 10 mL acetonitrile. The mixture to 25 mL volumetric flask was transferred with the aid of three portions of 2 mL acetonitrile. A mixture of water and acetonitrile (5:3) was prepared for mobile phase. Chromatograph equal volumes of the assay preparation and the standard preparation at 254 nm. Three repeated injections of the standard preparation were shown as a resolution factor of not less than 2 between the peaks for norethindrone and FLU and a relative standard deviation of not more than 1.5%.

2.10. Assay of Preservative (Parabens)

Approximately 5 g of cream was weighed in a 500 mL glass tube with screw cap. Five mL sulfuric acid solution (4/9) and 50 mL ethanol/water mixture (4/10) were transferred to the tubes. The tube was kept for five minutes in a water bath at 60 ± 1°C to facilitate the extraction of the preservatives into the ethanol phase. The extraction tube was rinsed with an ethanol/water mixture (4/10). A mixture of water/acetonitrile mixture (4/10) was prepared for mobile phase. Chromatograph equal volumes of the assay preparation and the standard preparation was screened at 254 nm. Methylparaben must be between 0.144% (NLT) and 0.216% (NMT). Propylparaben must be in the range of 0.016% and 0.024%.

2.11. Microbial Assessments

Ten g of cream was mixed with 3 g of sterile tween 80, then 24 g of 10% tween 60 water solution was added to TSB medium to diagnose the bacteria (Mixture A). Plates having 25 - 250 colonies were counted and the average number of CFU/g was calculated. Two dilutions (10¹, 10²) with the TSB culture medium were prepared and 1 - 2 mL of premixed dilution was added to 2 sterile plates containing 10 - 20 mL of TSA culture medium, and incubated for 24 - 48 hours at 30 - 35°C (Pure plate). One mL of 0.1 dilution for fungal counting allowed the agar to solidify at room temperature, then for 5 - 7 days incubated at 20 - 25°C. Colonies
were counted after the incubation and all operations were carried out in duplicate. All microbial counts were transformed into logarithms of the number of colony-forming units (log 10 CFU/g). Counting was passed in plates that contain less than 100 colonies for microbial count (TAMC) and contain less than 10 colonies for fungal count (TYMC) (18, 19).

2.13. *Staphylococcus aureus* Count

Colonies of *S. aureus* were counted after 24 - 48 hours of incubation at 30 - 35°C.

2.14. Yeast and Mold Counts

After the incubation period, the presence of white colonies and their confirmation with differential tests such as tuberculosis was confirmed by *Candida albicans*.

2.15. Preservative Effectiveness Test

A standardized inoculum containing 3 bacteria (*S. aureus*, *E. coli*, and *P. aeruginosa*), a yeast (*C. albicans*) and a mold (*Aspergillus niger*) was mixed with the liquid product. The mixture containing 0.05% of polysorbate 80, and add sufficient sterile buffer phosphate (pH: 7.2) to gain a count of about 1 × 10^8 CFU per mL. The absorption of this inoculum was read at 580 nm. Ten g of cream was added to 90 cc buffer phosphate (Mixture B) and 10^2 and 10^3 dilutions of the mixture B was made. Then, 1 mL of 10^2 and 10^3 dilutions was added to a sterile plate containing 15 - 20 mL of TSA medium, which cooled to 45°C, and incubated for 24 - 48 hours at 30 - 35°C. The fungal method was similar to the above except that the SDA culture with 10^2 dilution was incubated for 5 - 7 days at 20 - 25°C. Positive and negative controls were used during the experiment. Microbiological tests were repeated for formulations at 25°C after 14 and 28 days of the preparation (19).

2.16. In Vitro Release Studies

*In vitro* release of HQ, TRE, and FLU from formulation was evaluated by using cellulose nitrate membrane (0.45 micron). The membrane was mounted on modified Franz diffusion cells with a surface of 12.56 cm^2 and a receptor volume of 34 mL. The receptor compartment was nonstop stirred at 100 RPM using the magnetic stirrer. The receptor phases for HQ, TRE, and FLU were methanol, isopropyl alcohol, and acetonitrile, respectively. Then, 0.5 g of the formulations was applied with a spatula in the donor compartment and the diffusion cells protected with an aluminum foil to avoid light exposure. The temperature was maintained at 37.0 ± 1°C by a water jacket. Sampling was done at 30, 60, 120, 180, and 240 minutes, and at each point, 3 mL aliquots were drawn from the receiver section. After that, an equivalent volume of suitable receptor fluid was put back to the receiver compartment to save sinking conditions. The profile of release was then designed. Samples were analyzed with a UV spectrophotometer (Wavelengths of 293 nm for HQ, 352 nm for TRE and 254 nm for FLU) (20, 21).

### 3. Results and Discussion

An effective treatment for epidermal melasma is a combination of HQ, a steroid, and TRE. The combination strongly inhibits the production of melanin without the harmful damage of melanocytes. Kligman and Willis recommended a preparation containing HQ 5%, TRE 0.1%, and dexamethasone 21-acetate 0.1%, to be applied daily for 5 to 7 weeks, and it was found to be effective in the treatment of melasma. Despite its effectiveness, this preparation contains high concentrations of TRE and HQ, and holds dexamethasone, which is a potent fluorinated steroid. Dexamethasone and betamethasone (steroids 9-α fluorine) have no advantage over FLU. Indeed, FLU (acetone derivatives from fluorine steroids) is superior in the treatment of hyperpigmentation. In FLU, halogenation in the 9-α position increases the strength by improving the target cell activity and reducing the conversion to inactive metabolites. Also, covering or removing the hydrophilic part in the position of 17 dihydroxy acetonide or 16-α-hydroxy, increases the lipophilicity of the molecule and thus increases the penetration of stratum corneum (22, 23).

A cream containing HQ 4%, TRE 0.05%, and FLU 0.01% applied daily for 8 weeks was more effective than other formulations (5). In this formula, lowering the concentrations of the components reduced the frequency of irritancy. Because of HQ and TRE, this triple compound cream susceptible to instability (e.g. phase separation, change color, and decrease of assay). In this study, we prepared a stable generic pharmaceutical product with an appropriate analysis method (5, 24).

The freshly prepared cream had a yellow light color and homogenous texture. In order to find the emulsion type, the dilution test by distilled water depicted O/W structure. In addition conductivity assessment of this formulation by conductivity meter showed an electrical conductivity of 0.41 milliseconds which confirmed O/W system. As the control paraffin showed zero electrical conductivity. Finally, microscopic images demonstrated the dispersion of oil droplets in the aqueous media (Figure 1).
Our triple cream was exposed to the stability study as per ICH guidelines for a period of six months. It was checked at 25 ± 2°C/60 ± 5%. During the period of storage, the formulation showed no variation in color, odor, pH, and phase separation (Table 1). As shown in Table 1, the mean pH at the initial, 1st, 2nd, 3rd, and 6th months was 5.2, 5.4, 5.7, 5.6, and 5.7, respectively. The viscosity at the initial, 1st, 2nd, 3rd, and 6th months was 13325, 14429, 15670, 17688, and 17148 cp one by one. Finally, the density of fresh cream was 0.99174, and at 1st, 2nd, 3rd, and 6th months was 0.9952, 0.9960, 0.99 and 0.994, respectively. A microbial assessment at the initial, 1st, 2nd, 3rd, and 6th months was 7, 0, 0, 2, and 0 colonies, and fungi counts was 1, 0, 0, 0, and 0 colonies, respectively, indicating that antimicrobial activity has been adequate and efficient.

The assay of active pharmaceutical ingredients in the formulation was desirable. As shown in Figure 2, in HPLC results, the retention time of FLU was in 3.7 minutes, the retention time of Norethindrone (standard of FLU) was in 6.5 minutes, the retention time of TRE was in 1.2 minutes, the retention time of methylparaben and propylparaben was in 2.5 and 4.5 minutes, respectively. Therefore, as shown in Table 1, the percent of assay of HQ, TRE, and FLU at the initial, 1st, 2nd, 3rd, and 6th months were found in range. The percent of assay of HQ was 99.80, 99.71, 99.15, 99.50, and 99.25 at initial, 1st, 2nd, 3rd, and 6th months, respectively. The percent of the assay of TRE was 98.52, 98.23, 98.91, 98.00, and 98.15 at 1st, 2nd, 3rd, and 6th months, respectively. The percent of assay of FLU was 106.4, 105.6, 104.75, 102.89 and 103.26 at 1st, 2nd, 3rd, and 6th months, respectively. In addition, preservatives were at the desirable antimicrobial activity range. The percent of assay of methylparaben was 0.112, 0.108, 0.107, 0.106, and 0.109 at the initial, 1st, 2nd, 3rd, and 6th months, respectively. The percent of assay of propylparaben was 0.013, 0.010, 0.012, 0.010 and 0.011 at the initial, 1st, 2nd, 3rd, and 6th months, respectively.

In vitro diffusion studies reveals that our formulation showed a proper drug permeation into cellulose nitrate membrane and good release of all actives within 120 minutes, which may reflect a reliable therapeutic effect of formulation. As depicted in Figure 3, at the time of 90 minutes, FLU was 100% released from the cream basement. The HQ was reached with the maximum release of 71% at 40 minutes and TRE was shown a sustained release during 100 minutes to obtain nearly 100% release. Many commercial TRE topical formulations deliver all TRE in half an hour, which causes skin irritations. In this triple cream, slow and constant release of TRE in the first 60 minutes, in combination with the fast release of FLU as an anti-inflammatory agent, suggests minimum skin erythema, peeling, and dryness.

4. Conclusion

Melasma treatment can be challenging and require long-term treatment with topical agents. Melasma may be cured using monotherapy or combination therapy. These therapies, joined with regular application of a UVA and UVB blocking sunscreen to avoid UV-induced recurrence, are the most common therapies for epidermal melasma. A triple combination of HQ, TRE, and FLU has been counseled to be the first-line topical treatment for this pigmenitary condition. In this study, for the first time in Iran, we loaded three sensitive active agents in one emulsion without any phase separation, oxidation, and active’s potency loss. We recommend this new depigmenting combination as a generic pharmaceutical cream would be useful for melasma as well as other indications such as after dermal-esthetic procedures.

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Figure 2. HPLC assay of A, TRE; the retention time of this substance is in 1.2 minute; B, FLU; retention time of this substance is in 3.7 minutes; C, Norethindrone (standard of FLU), the retention time of this substance is in 6.5 minutes; D, both FLU and Norethindrone in one injection; E, methyl paraben and propyl paraben, the retention time of methylparaben is in 2.5 minutes and propyl paraben is in 4.5 minutes.

Figure 3. In vitro penetration and cumulative drug release of A, FLU; B, HQ; C, TRE; permeated through cellulose nitrate membrane.
### Table 1. Stability Evaluation and Antimicrobial Test Results

| Test | Period of Storage | Standard |
|------|-------------------|----------|
|      | Initial | 1st M | 2nd M | 3rd M | 6th M | Yellow light cream, homogenous cream mass |
|      | Complies | Complies | Complies | Complies | Complies |
| **Description** | | | | | |
| Odor | Complies | Complies | Complies | Complies | Complies | Characteristic odor |
| pH | 5.2 | 5.4 | 5.7 | 5.6 | 5.7 | NLT 5.00; NMT 6.00 |
| Viscosity (cp) | 13325 | 14429 | 15670 | 17648 | 17948 | NLT 10000; NMT 18000 |
| Density (g/cm³) | 0.99174 | 0.9952 | 0.9960 | 0.99 | 0.994 | NLT 0.980; NMT 0.999 |
| Assay (g/100 g cream) | | | | | |
| HQ | 99.80 | 99.71 | 99.15 | 99.50 | 99.25 | 94 - 106 % |
| TRE | 98.52 | 98.23 | 98.91 | 98.00 | 98.15 | 90 - 120% |
| FLU | 106.4 | 105.6 | 104.75 | 102.89 | 103.26 | 90 - 110% |
| Methyl paraben | 0.112 | 0.108 | 0.107 | 0.106 | 0.109 | 0.05 - 0.2% |
| Propyl paraben | 0.013 | 0.010 | 0.012 | 0.010 | 0.011 | 0.01 - 0.04% |
| Identification | Approved | Approved | Approved | Approved | Approved | The chromatograms of assay preparation and standard preparation exhibit the same major peak for cream |
| Microbial enumeration & tester for specific microorganism | | | | | |
| Total bacterial count (CFU/g) | 7 | 0 | 0 | 2 | 0 | NMT 100 |
| Total fungi & yeast count (CFU/g) | 1 | 0 | 0 | 0 | 0 | NMT 10 |
| *P. aeruginosa* | Neg. | Neg. | Neg. | Neg. | Neg. | Must be negative |
| *S. aurus* | Neg. | Neg. | Neg. | Neg. | Neg. | Must be negative |
| *E. coli* | Neg. | Neg. | Neg. | Neg. | Neg. | Must be negative |
| Antimicrobial effectiveness test | Ok | Ok | Ok | Ok | Ok | Bacteria: Not less than 2 log reduction from the initial count of 14 days, and no increase from 14 days’ count at 28 days. Yeast and mold: No increase from the initial calculated count of 14 and 28 days. |

### Footnotes

**Conflict of Interests:** The authors declare no conflict of interest.

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