Protective Effects of a Novel Preparation Consists of Concentrated Dead Sea Water and Natural Plants Extracts against Skin Photo-Damage

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ABSTRACT

Background: Exposure to solar UV rays has deleterious effects on skin appearance through physiological and structural alterations that eventually lead to skin photo-damage. Aims: To test the photo-protective effect of a novel preparation, Dead Sea Osmoter Concentrate (DSOC), containing high concentration of Dead Sea water. Methods: Human skin organ culture was used as a model to assess the biological effects of UVB irradiation and the protective effect of topical application of DSOC preparation. Skin pieces were analyzed for mitochondrial activity by Alamar blue (resazurin) assay, for apoptosis by caspase 3 assay, for detoxification by proteasome 20S sub-unit activity and for skin hydration by aquaporin 3 (AQP3) membrane channels expression. Human subjects were tested to evaluate the effect of DSOC Serum, a topical formula for facial skin, on skin hydration by a corneometer. Results: UVB irradiation induced cell apoptosis in the epidermis of skin organ cultures and decreased their proteasome activity and AQP3 expression. Topical application of DSOC significantly attenuated all these effects. In human subjects, an elevation in the intense skin moisture, which was higher relatively to subjects’ regular moisturizer, was observed. Moreover, DSOC Serum boosted the hydration effect together with regular moisturizer. Conclusions: The results clearly demonstrate that DSOC preparation can significantly improve the skin capability in coping with UVB rays stress in different cellular functions such as anti-apoptotic properties as well as detoxification and hydration recovery that can attenuate biological effects of skin photo-damage. Topical application of DSOC Serum had contributed to skin appearance by a strong hydration impact, also as a booster.

KEYWORDS

Photo-Damage; Skin Dryness; Skin Barrier Function; Skin Detoxification; Dead Sea Minerals

1. Introduction

Skin, as the outermost organ is exposed extensively to UV radiation-derived from sunlight, that can stimulate photochemical and photo-biological reactions, which eventually leads to tissue damage and could cause skin cancer [1,2]. Chronic exposure to UV radiation is involved with photo-damage and accelerates skin aging characterized by dry and rough surface, reduction of elasticity, wrinkles’ formation and hyper-pigmentation [3]. Skin exposure to UVB rays (280 - 320 nm) is related to photo-damage associated mainly with epidermis layers but it can also affect the dermal layer [4]. Since UVB photo-damage is an important component of UV damage, a protection against UVB photo-damage is very important from both skin care and dermatology points of view. Epidermis is one of the paramount target organs to en-
vironmental oxidative damage due to its location, as our body outermost layer, where it is always in contact with oxygenic air and occasionally exposed to UVB (mainly solar) light. Over-production of reactive oxygen species (ROS) could result in structural and functional alterations of cutaneous proteins resulted in loss of enzymatic activity, altered cellular functions such as energy production, interference with the creation of membrane potentials, and changes in the type and level of cellular proteins. Protein oxidation products are usually aldehydes, keto compounds, and carbonyls [5].

Additional major characteristic of photo-damaged skins is its dehydration status. Water is essential for vital skin functioning and particularly its outer layer, the stratum corneum (SC). Environmental daily insults, including sun exposure, hot dry air, and winds might lower the SC water content, causing improper desquamation and the appearance of dry, flaky skin [6]. Aquaporin 3 (AQP3), a member of the family of aquaglyceroporins, is a small, hydrophobic, integral membrane protein that act primarily as water-selective pores, thus contributes to balance the water level and to water loss prevention [7]. AQP3 provides a short circuit for water between the base of the epidermis and the SC, maintains constant water content, and thus susceptible to pre-mature skin aging. Sunscreen application supplies only limited defense against the long-term consequences of UV exposure, therefore, additional treatment strategies should be considered for enhance skin protection. Various natural components can contribute to non-sunscreen solar protection and their activity can assist in skin protection by intervention in metabolic pathways involved in UV damages.

Dead Sea (DS) water, salts, minerals and mud are well known for their therapeutic activities and are used in treating a variety of skin conditions as well as for their cosmetic benefits [9,10]. Osmoter™, a natural commercial composition of DS water was shown to reduce skin senescence markers, to attenuate UVB damage markers [11] and to improve skin smoothness [12]. Inclusion of extracts isolated directly from plants and other natural sources for therapeutic and cosmetic purposes can contribute and improve solar protection.

Dead Sea Osmoter Concentrate (DSOC) preparation (US patent application pending 61/812,946), formulated by AHAVA Dead Sea Laboratories LTD, is a unique combination of natural actives consisting of concentrated Dead Sea Minerals extract (Osmoter™), at a relatively high concentration, with Green tea extract, Grape seed extract, Olive leaf extract and Benzotriazolyl Dodecyl p-Cresol in water. These extracts were selected based on bio-activity measured in a screening process of a list of candidate actives we had performed ex vivo on human skin organ cultures with and without exposure to UVB irradiation (data is not shown).

In this study, the capabilities of DSOC to protect human skin against UVB-induced biological damaging impacts were investigated on ex-vivo human skin model and in-vivo clinically on human volunteers.

2. Methods

2.1. Treatment Preparations

All preparations used in this study were manufactured by AHAVA Dead Sea Laboratories, Ltd., Israel.

2.2. Dead Sea Osmoter Concentrate™ Preparation

Dead Sea Osmoter Concentrate™ (DSOC) preparation: Osmoter™ solution, Benzotriazolyl Dodecyl p-Cresol (Tinogard TL), green tea extract (Greenselect), grape seed extract (Leucoselect) and olive leaf extract (Eurol BT) in water.

Osmoter™: The major constitutes of DS water are the following ions: Mg²⁺ (92,500 mg/l), Ca²⁺ (38,000 mg/l), K⁺ (1400 mg/l), Na⁺ (2000 mg/ml), Sr²⁺ (800 mg/l), Cl⁻ (345,000 mg/l), Br⁻ (11,500 mg/l).

2.3. DSOC Serum Preparation

DSOC preparation was further incorporated into topical preparation in a form of a serum (DSOC Serum). The various additional ingredients comprised within the DSOC Serum are as follows: Aqua (Mineral Spring Water), PVP, Maris Aqua (Dead Sea Water), Cyclomethicone, Dimethicone, Butylene Glycol & Sigesbeckia Orientalis Extract & Rabdosia Rubescens Extract & Aqua (Water), Caprylyl Methicone & PEG-12 Dimethicone/PPG-20 Crosspolymer, Aloe Barbadensis Leaf Juice, Glycérine, Isodecyl Isononanoate, Diisobutyl Adipate, Dimethicone & Dimethiconol, Phenoxethanol & Aqua (Water) & Biosaccharide Gum-1, Sodium Lactate, Trehalose, Butylene Glycol & Phenoxethanol & Laminaria Digitata Extract, Citric Acid & Glycerin & Lactic Acid & Parfum (Fragrance) & Phenoxethanol & Aqua (Water) & Ascorbic Acid & Citrus Reticulata Fruit Extract & Citrus Aurantium Amara Fruit Extract, Bisabolol, Squalene (Phytosqualene), Argania Spinosa Kernel Oil, Parfum (Fragrance), Hydroxyethyl Acrylate/Sodium Acryloyldimethyl Sodium Hyaluronate, Benzotriazolyl Dodecyl p-Cresol, Helianthus Annuus (Sunflower) Seed Oil & Aqua (Water) & Ascorbic Acid & Melissa Officinalis Leaf Extract & Polyglyceryl-3 polyricinoleate, Tris(tetra-methyl hydroxyl piperidinol) citrate, Sodium Benzotriazolyl
Butylphenol Sulfonate, Glucosyl Hesperidin, Aqua (Water) & Olea Europaea (Olive) Leaf Extract & Fructose, Pentaerythrityl Tetra-di-t-butyl Hydroxyhydrocinn, Vitis Vinifera (Grape) Seed Extract & Phosphatidylcholine, Camellia Sinensis (Green Tea) Leaf Extract & Phosphatidylcholine.

2.4. Human Skin Organ Culture Model for Biological Tests

Skin fragments were obtained with informed consent from 20 - 60 years old, healthy women, who underwent breast or abdomen reduction. Samples were cut into pieces of 5 × 5 mm. Each experiment consisted of 4 replications for each treatment. The examined preparations were applied as a thin layer on the epidermis of each skin piece (2.5 μl/piece). The skin pieces were incubated in 12-wells culture plates filled with DMEM culture media (Dulbecco’s Modified Eagle’s Medium, Biological Industries BeitHaemek, Israel); (37°C; 5% CO2) so that the epidermis was exposed to the air while the dermis was submerged in the culture media. After 24 hr of incubation, the preparations were wiped off from the skin on a sterile filter paper, the skin was washed in PBS and irradiated by a UVB lamp (VL-6.Mlamp, emission spectrum 280 - 350 nm, emission peak 312 nm, filter size 145 × 48 mm, Vilber Lourmat, Torcy, France.). The irradiation intensity was 2 mJ/cm²/sec. Irradiation time was calculated in order to supply 250 ml/cm². Freshly preparations were applied on the skin pieces and they were further incubated in culture plates containing replenished medium.

At the end of the post-irradiation period (which depended on the examined parameter), the skin was incubated for 1 min in phosphate buffered saline (PBS) at 56°C and the epidermis was separated from the dermis with a scalpel. The remaining culture medium was collected, frozen and kept at −70°C until use.

2.5. Apoptosis Determination by Caspase 3 Activity Assay

Apoptosis determination by caspase 3 assay was carried out 24 hr following UVB irradiation. For Caspase-3 activity measurement, the collected epidermis pieces were placed into wells of a 96-well plate, each filled by 125 μl of Caspase-3 specific substrate solution (10 μM Ac-DEVD-AMC, Merck, Darmstadt, Germany) with 0.02% Triton X-100 (J. T. Baker, Phillipsburg, NJ, USA) and 10 mM DTT (TCI, Tokio, Japan) (34). The enzyme’s fluorescent product emission level (355/460 nm) was measured at 37°C at 2-minute intervals using fluorescence plate reader Fluostar-BMG spectrofluorimeter (Offenburg, Germany) for 40 min. Apoptosis level was later deduced from the slope value of enzymatic activity in the linear range.

Osmoter™ solutions in water at a concentration range of 0.8% and 4% were tested on ex vivo human skin organ cultures in order to determine the relevant Osmoter™ concentration in DSOC.

2.6. Viability Measurements by Mitochondrial Assay

Skin viability was determined using Alamar blue test using resazurin reagent, which is a redox indicator that both fluoresces and changes color in response to chemical reduction of the growth medium resulting from cell growth. Fresh growth medium containing 0.01 mg/ml resazurin (Sigma Aldrich, Israel) was added to cultures at the end of the test period and incubated for 2 hrs. Fluorescence at 540 nm (excitation)/590 nm (emission) proportional to mitochondrial activity was measured on a BioTek Synergy 2 microplate reader.

2.7. Proteasome Activity Determination

Proteasome activity of 20S sub-unit was measured following DSOC preparation application 72 hr post irradiation on epidermis samples as adapted for Bregere et al. [13]. For testing its chymotrypsin-like and trypsin-like activity, epidermis samples were peeled and incubated in 125 μl PBS containing LLVY-AMC as a substrate, with 0.02% Triton X-100 and 10 mM DTT, at 37°C in a 96-well plate Fluorescence of the released coumarin derivative was measured at 360/460 nm, using a Fluostar-BMG spectrofluorimeter. Activity was measured by the fluorescence-versus-time slope, calculated over 30 min in the linear range.

2.8. Aquaporin 3 (AQP3) Levels Determination

The degree of Aquaporin protein expression in epidermis was determined following DSOC preparation application 48 hr post irradiation. Prior to that, AQP3 expression was tested on epidermis 48 hr following application of Osmoter™ in different concentrations without irradiation. AQP3 levels were determined by dot-blot analysis. Each separated epidermis sheet was placed in 500 μl of sample buffer (Tris 0.25 M pH 6.8, SDS 1%, Glycerol 5%, β-mercaptoethanol 5%), boiled for 5 minutes and kept at 20°C until use. Protein concentration in boiled samples was determined. Briefly, 3 μl of each sample were spotted onto Whatman paper 3 MM, air-dried, rinsed with methanol and stained with Coomassie stain. After comparison to BSA standard curve, all the samples were diluted to 0.33 g/ml in sample buffer. For immuno analysis, 3 μl of tthew samples (1 μg of protein), in 4 replicates were deposited directly onto nitrocellulose membrane.
(dot blots). The membrane was processed as using anti-
gen-specific primary antibody (ab15117, Abcam Cam-
bidge, UK) and horseradish peroxidase-conjugated, anti-
rabbit secondary antibody (Jackson Immuno Research, West Grove, PA, USA). Chemoluminescence was de-
tected using a substrate cocktail composed of: 1.25 mM luminol, 0.2 M p-coumaric acid and 0.009% H₂O₂ in 100 mM Tris/HCl pH 8.8, in a Fujifilm LAS 3000 instrument (Fujifilm, Tokyo, Japan). Densitometric analysis was per-
formed using the NIH Image J program.

2.9. Clinical Measurements for Skin Hydration

Measurements of skin hydration were performed by Dermatest GmbH medical research clinic, in Munster, Germany. The study was carried out on 35 female sub-
jects aged 18 - 74 years on the forearm.

No patient was enrolled without his or her signed and
witnessed informed consent. The study was approved by
the Institutional and Ministry of Health Ethics Commit-
tee.

The measurements were performed over a defined pe-
oriod of time—(0.5-hr, 8-hr, 24-hr), after a single applica-
tion to 4 different application areas in the flexor aspects of
the forearms. In the first area only the DSOC Serum
was applied, in the second area only the Regular Moistu-izer of the subjects, in the third area the DSOC S erum
plus the Regular Moisturizer. The last area served as
control area, nothing was applied.

The subjects were instructed to apply only the cosmetic
preparation being tested on the relevant pre-deter-
rmined skin areas during the application period. The test
subjects were acclimatized for 45 minutes at a tempera-
ture of 22°C and 60% relative humidity. Skin values
were measured at three different places within the re-
spective testing areas.

Electrical capacitance was measured with a capacitan-
city meter (Corneometer CM 825, Courage & Khazaka, CK). The probe head consisting of
a condenser was applied on the skin surface at constant
pressure. The range values of each measurement were
recorded and compared to the other measurements.

The recorded values were averaged. For this experi-
ment, the untreated skin close to the test area was used as
the control measurement area. Measurements were taken
before the application and 30 minutes, 8 hours and 24
hours after a single application.

The results represent the average percentage moisture
change caused by the application with reference to start-
ing value.

Inclusion criteria: subjects with a lack of skin tone
homogeneity/slight discolorations and with dehydrated
skin in the test area aged 18 years and older.

Exclusion criteria: severe or chronic skin inflamma-
tion; serious inner or chronic diseases; intake of drugs
that possibly can interfere with skin reactions (Glucocor-
ticoids, antiallergics, topical immuno modulator, etc.);
application of pharmaceutical products and skin care
products with active ingredients until 7 - 10 days before
testing; severe allergies or occurred severe side effects
after usage of cosmetic products; sunbath or usage of
tanning bed during the study period; known cancer;
pregnancy or lactation period.

2.10. Adverse Events and Tolerance

Before the start of the application test all participants
were determined to have healthy skin in the test area. No pathological skin disorder was detected. During the study:
No complaint of any pathological skin disorder was re-
ported during the course of this application test. Interru-
tions of the application test and/or medical intervention
were not necessary.

After the end of the application test: During the final
dermatological examination after the end of the study,
none of the 35 participants showed development of any
pathological skin disorder in the test area. The mentioned
product was well-tolerated and did not lead to any un-
wanted skin reaction.

2.11. Data Analysis

Average values are given with standard error of the mean
(SEM). Differences between average values were tested
for significance using the Two-tailed Student t-test for ex
vivo experiments and Dunnett test for in vivo
experiments and considered significant for \( p \leq 0.05 \).

Each ex vivo experiment was performed at least in triplicate.

3. Results

3.1. Ex-Vivo Results

The efficacy of topical application of DSOC preparation:
Skin specimens were applied with DSOC preparation
and UVB-irradiated. Skin biomarkers for apoptosis, viability,
detoxification and hydration were measured in skin spec-
cimens with and without UVB irradiation stress.

The significant enhancement in pro-apoptotic caspase
3 enzyme activity-induced by UVB irradiation, was
markedly decreased when skin was pre-treated topically
with DSOC preparation: 34% compared with non-treated
samples \( (p < 0.05) \) (Figure 1). The application of DSOC
preparation did not induce apoptosis in non-irradiated
samples.

Alamar blue (resazurin) mitochondrial activity assay
was used for analyzing skin viability. Topical application
of DSOC preparation did not impede mitochondrial ac-
tivity in both non-irradiated and irradiated samples.
Moreover, pretreatment by DSOC preparation enhanced

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mitochondrial activity in irradiated samples by 75% (p < 0.01) (Figure 2).

The proteasome is the major intracellular proteolytic system implicated in the deletion and recycle abnormal and oxidized proteins. In human epidermal cells, previous studies have evidenced that proteasome function is decreased during aging as well as upon UV irradiation. Proteasome activity was used to indicate the rate of degradation of unwanted or damaged proteins in the epidermis. Results showed that UVB irradiation in untreated samples led to a significant decrease of 40% (p < 0.05) in 20S sub-unit activity (Figure 3). Pre-treatment with DSOC preparation prevented this decrease.

AQP3, a membranal protein acting as water/glycerol transporter, is an important contributor for skin hydration.

UVB irradiation decreased AQP3 protein levels in epidermis of untreated samples by 17% (p < 0.05) (Figure 4). Pre-treatment with DSOC preparation markedly abolished this decrease: AQP3 level in irradiated treated samples was 88% higher than in irradiated untreated ones (p < 0.05). Also, an increase in AQP3 expression by 44% (p = 0.05) was observed in non-irradiated skin pre-treated with DSOC compared to non-irradiated control.

### 3.2. In-Vivo Clinical Results

Skin hydration was evaluated before and after topical application of the following treatments: DSOC Serum, Regular Moisturizer and DSOC Serum together with topical application of regular moisturizer. A significant elevation in skin hydration was presented 0.5 hr and 8 hr after all treatments application compared to the control area: 105% for DSOC Serum (p < 0.01), 90% for Regular Moisturizer (p < 0.01) and 136% for DSOC Serum + Regular Moisturizer (p < 0.01) after 0.5 hr; 50% for DSOC Serum (p < 0.01), 41% for Regular Moisturizer (p < 0.01) and 51% for DSOC Serum + Regular Moisturizer (P < 0.01) after 8 hrs. (Figure 5). 24 hr after treatments application, a significant elevation in skin hydration was observed only after DSOC Serum and DSOC Serum + Regular Moisturizer application compared to control area: 17% for DSOC Serum (p < 0.01) and 21% for DSOC Serum + Regular Moisturizer (p < 0.01).

### 4. Discussion

Skin exposure to Ultraviolet (UV) radiation plays an important role in the pathogenesis of skin aging and skin cancers. Epidermis, the outermost skin layer, is primarily affected by UV Brays, which can lead to photo-damaged phenotype due to its involvement in molecular oxidation,
inflammation and dryness [14]. Hence, a topical application of protecting preparations, possessing biological activities, such as recycling damaged oxidized substances (detoxification), anti-inflammatory properties and at the same time maintaining skin moisture, may be able to attenuate the process of skin photo-damage.

This study is focused on a Dead Sea Osmoter™ Concentrate (DSOC), a novel preparation consisting of Osmoter™—a special mixture of DS water extract highly concentrated, Green tea extract, Grape seed extract, Olive leaf extract and Benzotriazolyl Dodecyl p-Cresol in water.

The therapeutic properties of DS water are well established and are related to the unique mineral composition, mainly dissolved salts of magnesium, calcium, sodium, potassium, zinc and strontium. Many DS based compositions have been used to also alleviate skin disorders symptoms. Studies have investigated the role and mechanisms of the DS minerals on skin [11]. Some studies have focused on isolating the effect of particular minerals and some of the semetallic elements from DS water have been proven to be involved in skin metabolism: zinc ions were reported to enhance epidermal proliferation and wound healing [15]. Magnesium and calcium ions are known as skin barrier recovery accelerators. DS water has been reported to stimulate cellular proliferation and mitochondrial activity, to decrease the expression of aging bio-markers and to diminish apoptotic damage after skin exposure to UVB irradiation [11]. In this study, we further examine the effect of DS minerals in association with plant extracts on essential skin functions and with regards to UVB irradiation, a main component of skin aging.

Green tea (Camellia Sinensis) leaf extract is rich in poly-phenols. The most abundant and biologically active poly-phenol is the catechinpigallocatechin-3-gallate (EGCG), which is a potent antioxidant. Many studies have shown that topical treatments or oral consumption of green tea or its poly-phenols can inhibit skin tumorigenesis, induced by chemical carcinogens or ultraviolet radiation. Recent Study has shown that Green tea leaf extract also possesses anti-inflammatory activity [16].

Grape (Vitis Vinifera) seed extract is rich in pro-anthocyanidins, potent antioxidants reported to possess more free radical scavenging activities than vitamins C and E [17]. Grape seed extract is also known to protect skin against UVB irradiation through inhibiting the depletion of antioxidant defense components induced by UVB [18].

Olive (Olea Europaea) leaves have long been used in folk medicine in Mediterranean countries. Oleuropein, the main polyphenolic compound in Olive leaf extract, and its derivatives have tested for a variety of biochemi-
cumulation of oxidatively modified proteins. Proteasome, the major intracellular proteolytic system implicated in the removal of abnormal and oxidized protein, has a pivotal role in cellular detoxification. Previous studies in human epidermal cells have shown that proteasome activity is decreased during senescence as well as upon UV irradiation [24]. It has been shown that keratinocytes UVA and UVB irradiation led to a decrease in proteasome peptidase activity. 20S sub-unit was found to be very sensitive in its active form to UVA- and UVB-irradiation mainly affecting the peptidylglutamyl-peptide hydrolase activity [24]. In this study, proteasome 20S sub-unichymotrypsin-like activity and trypsine-like activity indeed decreased in skin organ cultures following exposure to UVB irradiation (Figure 3). Pre-treatment with DSOC preparation abolished this decrease and thus, assisted in maintaining protein turnover, which is essential to preserve cell function. This finding can be related to DSOC preparation neutralizing ROS formation. A study published in 2011 article by Koziel et al. suggests a functional interplay between mitochondrial activity and proteasome activity in skin aging, proposing that both systems can be associated in skin aging [25]. Since DSOC is shown to induce mitochondrial activity, indeed, this correlation has been found between the two functions with regards to UVB photoaging.

One major result of photo-damage is the dehydration of the skin [8]. Aquaporin 3 (AQP3)—membrane-inserted water channel play an important role in the regulation of water permeability [7]. Indeed, pre-treatment with DSOC markedly enhanced AQP3 expression in epidermis (Figure 4).

AQP3 expression has been linked to various skin disorders. Based on a study showing the aquaporin elevation after MgSO4 administration in rat [26], it might be speculated that in the case of DS water, the rich content of magnesium salt could also contribute to elevation in AQP3 expression.

Previous studies have demonstrated that UV radiation down-regulates AQP3 expression in cultured skin keratinocytes, leading to a decrease in water permeability also due to the reduced glycerol transport through AQP3 [27].

AQP3 down-regulation can also be due to UV-induced ROS over production via activating the MAPK/ERK (mitogen-activated protein kinases/extracellular signal-regulated kinase) pathway as shown in cultured skin keratinocytes [28].

Here, we show that AQP3 protein expression is decreased following UVB induction in human skin organ model while pre-treatment with DSOC preparation has a protective effect against UVB by increasing AQP3 levels (Figure 4).

Based on its beneficial effects on human skin organ culture model in different modes of biological activities, DSOC preparation was formulated into a serum (DSOC Serum) for facial skin and was tested for skin hydration. Skin hydration is important for its health and functionality. A decrease in skin moisturization level is related to different biological processes and stressors (i.e., aging, UV radiation, autoimmune skin disorders, and harsh environmental climates) and might lead to the impairment in skin barrier function, enhancement in skin roughness, and decrease of skin elasticity and firmness[21,29].

DSOC Serum demonstrates intense moisturizing effects by significantly increase SC hydration level (Figure 5). The improvement in hydration was the higher relatively to subjects regular moisturizer. Furthermore, DSOC serum boosts the effect of hydration together with the regular moisturizer. These results from the clinical study may reflect in addition to moisture at the skin surface level via hygroscopic molecules and emollients, the positive effect at the cellular level of DSOC on AQP3. Additional studies can be performed to elucidate the contribution of other actives in the DSOC Serum to skin.

5. Conclusion

In conclusion, we showed that a selection of plant extracts together with DS water extract (DSOC preparation) can significantly assist the skin in coping with UVB rays stress via anti-apoptotic properties and by maintaining normal proteasome activity and AQP3 expression. For the first time, it was demonstrated that concentrated DS water combined with plant extracts affect directly on proteasome activity and AQP3 expression. Hence, DSOC
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Figure 5. Skin hydration between control forearm and tested hand forearm at different time points following application of DSOC Serum compared to other treatments. Skin hydration in the flexor aspect of the forearm was measured in 35 female volunteers using a corneometer as described at the methods section. The measurements were performed in defined periods after a single application in 4 different application areas for DSOC Serum, Regular Moisturizer of the subjects, DSOC Serum plus the Regular Moisturizer and control (non-applied) area. The relative improvement between each treatment and control area was calculated. Data are presented as Mean ± SEM for delta (%), i.e., percentage moisture change caused by the application, with reference to the starting value (before application). **p < 0.01.

protects skin from photo-damage simultaneously on different cellular functions.

DSOC preparation implemented in a cosmetic formula (DSOC Serum) was shown in a clinical study to increase significantly skin moisture and to also boost the moisturizing capacity of regular moisturizer and is thus, expected to improve facial skin appearance. Further studies are needed to complete the understanding of DS minerals on skin moisture. The impact of UVA is needed to fully examine the protective effect of DSOC of aquaporin and proteasomal function against photo-damage.

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