SHORT REPORT **Topical Application of M89PF Containing Vichy** Mineralising Water and Probiotic Fractions Prevents Cutaneous Damage Induced by Exposure to UV and O_3

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Purpose: Exposure of the skin to ultraviolet radiation (UV) or ozone (O₃) results in stressed skin, leading to the alteration of the skin physical barrier and defence functions. In this work, the preventive benefit of a dermocosmetic, M89PF, containing Vichy mineralising water, probiotic fractions, antioxidant vitamins and hyaluronic acid, in the alteration of skin physical barrier and skin defence functions after exposure to O₃ and UV, alone or combined, was assessed.

Methods: Untreated and treated (M89PF) skin explants were exposed to O_3 , to UV rays or to O_3 +UV. Immunofluorescence was performed for skin barrier, oxidative stress, and inflammatory markers after one and four days of exposure to the pollutants.

Results: M89PF significantly (p≤0.05) prevented the decrease of the expression level of different skin barrier markers, and significantly $(p \le 0.05)$ prevented the induction of OxInflammatory markers and inflammasome components by UV, O₃ or both combined.

Conclusion: M89PF prevents skin barrier damage, as well as oxidative stress and inflammatory markers induced by exposome factors, such as UV, O₃ or both combined.

Keywords: ultraviolet radiation, ozone, skin barrier, oxidative stress, inflammation

Introduction

The skin is the largest and most diverse organ of the human body with the epidermis being the first line of defence against the skin exposome which includes ultraviolet radiation (UV) and pollution.¹⁻⁵ The effect of pollutants, such as Ozone (O₃) and UV on human skin has been investigated over the past decades. Ferrara et al have studied the combination of UV and pollution, confirming their impact on both the skin barrier and skin defence system through oxidative stress and activation of inflammatory cytokines.⁶

UVB and UVA significantly impact the skin, causing sunburn and skin aging.^{7,8} O₃ interacts with lipids and proteins present in the stratum corneum, resulting in the production of reactive oxygen species (ROS), secondary mediators such as aldehydes (4-HydroxyNonenal, 4-HNE) and cyclooxygenase 2 (COX-2), and in the activation of inflammasome pathways, such as NLR Family Pyrin Domain Containing (NLRP1).⁹⁻¹⁵ This process causes oxidative damage throughout the skin and initiates inflammatory processes.^{11,16-20} A growing set of data demonstrated that different pollutants affect skin expression levels of the transcription factor arvl hydrocarbon receptor (AhR) and that its activation in skin cells parallels that of keratinocyte differentiation and inflammation, resulting in cutaneous damage, skin diseases and premature skin aging.^{21–23}

Moreover, O₃ and UV alter the expression of the stratum corneum cornified envelope components such as lipids, proteins (Keratin, Filaggrin, Involucrin), as well as tight junctions.^{15,24–31} As a result, skin is stressed, which leads to the alteration of the skin physical barrier and defence functions.^{32,33} Filaggrin (FLG), a marker of the late epidermal differentiation, plays a pivotal role in the formation and maintenance of the skin barrier, as well as in stratum corneum hydration. Involucrin (INV), a protein involved in the assembly of the cornified envelope, the structure formed in the outermost layers of the epidermis, provides a physical barrier against environmental stressors.³⁴ Finally, as mentioned before, the exposome induces the activation of the inflammasome, a cytosolic multiprotein complex involved in skin inflammatory diseases, upon exposure.^{31,35} Inflammasomes represent a group of protein complexes of the initiation of an inflammatory status. Ferrara et al described the role of inflammasome activation in different skin pathologies and its possible role as a molecular target in the prevention of pollution-induced skin damage.³⁶ Among the different proteins involved in the inflammasome complex, the apoptosis-associated speck-like protein containing a C-terminal caspase recruitment domain (ASC), can be considered a hallmark of its activation.³⁷

M89PF (Mineral 89 Probiotic Fractions, Laboratoires Vichy, France) contains Vichy mineralizing water (VMW), hyaluronic acid, niacinamide, tocopherol and probiotic fractions. It is hypoallergenic and contains no perfume. VMW originates from the French volcanic region and contains 15 minerals with a total mineral concentration of 5.2g/l.

This ex vivo study assessed the effect of O_3 and UV radiation alone or combined on the skin, as well as the preventive effect of M89PF.

Materials and Methods

Maintenance in Culture and Exposure to UV and O_3 of ex vivo Skin Explants

All human tissues were obtained from elective abdominoplasty with donor consent under Pearl IRB approval, in accordance with FDA 45 CFR 46.102 and 21 CFR 56.102 regulations (Pearl Pathways. Exemption Determination Submission. IRB Study Number: 21-TENB-101. Study Title: Collection, culture, and distribution of human abdominoplasty skin tissue). Written informed consent was obtained from all subjects or, if subjects were aged less than 18 years, from a parent and/or legal guardian. All donors were healthy volunteers. No identifying information beyond ethnicity, sex and age was provided.

Subcutaneous fat was trimmed from 12 mm punch biopsies, and samples were rinsed with Phosphatase Buffer Solution (PBS) 1X containing antibiotics/antimycotic. Biopsies were cultured in DMEM Media added with 10% FBS, 100 U/mL penicillin, and 100 μ g/mL streptomycin at 37 °C in 5% CO₂ in 6-well plates, with the upper part of the epidermis exposed to the outside environment. The day after, the media was changed, and M89PF was topically applied. After 24 hours, the media was replaced with fresh media, and biopsies were exposed to 200 mJ UVA/UVB light alone (Newport, Oriel1, Sol1ATM, 1600 W, Xenon Lamp, UVC & AM0 filters), to 0.25 ppm of O₃ alone for 2 hours, or to a combination of 200 mJ UVA/UVB and 0.25 ppm of O₃ for 2 hours. The UVA/UVB ratio used was 21/1 since the air mass filter AM 0 was adopted to adjust the light output to strictly match the solar spectrum when the sun is at a Zenith angle of 0. UV doses were precisely monitored using a radiometer ILT2400 Hand-Held Light Meter / Optometer (International Light Technologies, Inc., Peabody, MA, USA). Biopsies were pre-treated every day with M89PF, exposed to the pollutants, and collected 24 hours after the first exposure (DAY 1) and after four days (DAY 4) for the subsequent evaluations.

Haematoxylin and Eosin Staining

Skin explants were fixed in 10% NBF (neutral-buffered formalin) and embedded in paraffin. For histological observation, sections (4 µm thickness) were deparaffinized in xylene, and rehydrated in a series of alcohol gradients (100%, 90%, 80% and 70%). Sections were stained with Mayer's hematoxylin solution, washed in tap water, stained with aqueous Eosin Y solution (Sigma Aldrich, St. Louis, MO, USA) for 2–3 min, and dehydrated in a series of increasing concentrations of alcohols and xylene. Finally, sections were mounted onto glass using a toluene based mounting medium and observed under microscope (Nikon Microphot FXA microscope,Nikon Instruments, Amsterdam, Netherlands).

Immunofluorescence

Paraffin sections (4µm thickness) of skin explants were deparaffinized in xylene and rehydrated in alcohol gradients. After that, sections were blocked with 5% BSA in PBS 1X for 45 minutes at room temperature, and incubated overnight at 4°C with the following diluted (in PBS 1X, 2% BSA solution) primary antibodies: 4HNE (diluted 1:500; AB5605, Millipore Corporation, Burlington, MA, USA), AhR (diluted 1:500; 83,200, Cell Signaling, Danvers, MA, USA), COX-2 (diluted 1:400; NB100-868, Novus Biologicals, Littleton, CO, USA), Involucrin (diluted 1:50; sc-21748, Santa Cruz Biotechnology, Inc., Dallas, TX, USA), Filaggrin (diluted 1:50; sc-66192, Santa Cruz). After 24 hours, sections were washed three times with PBS 1X, incubated with specific fluorochrome-conjugated secondary antibodies (diluted 1:500 in PBS 1X, 2% BSA; Alexa Fluor 568 A11004 or Alexa Fluor 488 A11055) at RT, and nuclei were stained with DAPI solution (D1306, Invitrogen) for 2 minutes. Sections were again washed three times with PBS 1X, and mounted using PermaFluor mounting media (Thermo Fisher Scientific, Waltham, MA, USA). Images of sections were made using a Nikon Microphot FXA microscope, and images captured were quantified using ImageJ.

Statistical Analysis

All results were expressed as mean \pm SD values of three different images. For between-group comparisons, an analysis of variance (ANOVA), followed by Bonferroni's post-hoc test, which compares all pairs of columns under test with a consideration of p value ≤ 0.05 as the significant difference, was conducted. Data were analysed using GraphPad Prism 6.0 (GraphPad Software, Inc., La Jolla, CA) software.

Results

Effect on the Skin Structure

By Haematoxylin and Eosin staining of skin explants, no evident alterations of the skin structure were observed after exposure to pollutants compared to control conditions, with an intact structure of the epidermis presenting all skin layers and a contiguous stratum corneum, validating the assay methodology. No noticeable changes were observed within the dermis when compared to control samples, suggesting that the selected dose of pollutants did not damage skin biopsies. No significant differences were perceived between treated and untreated tissues, suggesting that treatment with M89PF does not affect the skin morphology during the experimental procedure at either DAY 1 or DAY 4 (Figure S1).

Effect on Hydration and Skin Barrier-Associated Proteins, and the Preventive Role of M89PF

A decrease of FLG levels in skin samples exposed to O_3 and UV was observed specifically at DAY 4 (Figure S2B). This depletion was prevented by M89PF. M89PF was efficient after the exposure to the combination of O_3 and UV at DAY 1 and DAY 4 (Figure S2A and B), with a significant increase of the FLG expression levels (p ≤ 0.05 ; M89PF+O₃ vs O₃=+27%, M89PF+UV vs UV=+43% and M89PF+O₃+UV vs O₃+UV=+94%).

As depicted in Figure 1, the immunofluorescence analysis at DAY 1 (Figure 1A) demonstrated a decrease of the INV expression levels after exposure to pollutants. The topical administration of M89PF prevented this effect not only at DAY 1 but also at DAY 4 (Figure 1B), as confirmed through statistical analysis of the protein quantification (Figure 1, right panels).

Pre-Treatment with M89PF Prevents the Pollutant-Induced Expression of Oxidative Stress Markers

Combined exposure to UV and O_3 induced the expression of oxidative markers. However, this effect was counteracted by the pre-treatment of skin tissue with M89PF.

A significant ($p\leq0.05$) increase of the 4-HNE expression levels were observed in untreated skin samples after exposure to UV (+75%) or to UV+ O₃ (+90%), compared to the negative control at DAY 1 (Figure 2A). No notable change was observed in M89PF-treated skin samples. Pre-treatment with M89PF prevented 4-HNE increase, observed after the exposure to O₃+UV (-35%; $p\leq0.05$ if compared to pollutant-exposed samples).



Figure I Preventive role of M89PF on the skin barrier-associated protein Involucrin. Involucrin expression levels in skin explants samples exposed to O_3 , UV, O_3 + UV and/ or pretreated topically with M89PF for 24h and analyzed at DAY I (**A**) or DAY 4 (**B**). Magnification 40X. Scale bar 100 μ m. Right panels shown the quantification of relative immunofluorescence staining for Involucrin after different treatments (DAY I; DAY4). Data are expressed as the mean of the three different subjects ± standard deviation. *p≤0.05 Ctrl vs Pollutant; [#]p≤0.05 M89PF + Pollutant vs Corresponding pollutant. One-way ANOVA followed by Bonferroni post hoc test.

After 4 days (Figure 2B), an increase of the 4-HNE expression levels for both O_3 and UV alone, as well as for the combination of the two pollutants, was observed; this difference was statistically significant (p \leq 0.05) for O_3 (+76%) and for UV (+72) exposure). At DAY 4, the preventive effect of M89PF was even more evident than at DAY 1.

The expression of the AhR levels in untreated skin samples significantly ($p\leq0.05$) increased after exposure to O₃ (+48%) or to UV+ O₃ (+61%) at DAY 1 (<u>Figure S3A</u>). In M89PF-treated UV+O₃-exposed skin samples, AhR expression was significantly reduced compared to untreated samples exposed to UV+ O₃ (-50%, $p\leq0.05$). The impact of pollutants was less pronounced, and differences to control samples were statistically insignificant after four days of exposure (Figure S3B). Nevertheless, in this case, M89PF also had a protective role against O₃ and UV exposure.

Pre-Treatment with M89PF Prevents the Pollutant-Induced Inflammatory Skin Response

The levels of COX-2 in untreated skin samples exposed to UV (+50%) and UV+O₃ (+31%) significantly ($p\leq0.05$) increased at DAY 1 (Figure 3A) compared to control tissues. Topical administration of M89PF resulted in a decrease of COX-2 expression levels in UV and UV+O₃ exposed samples (about -30%). Exposure to O₃ alone did not alter the COX-2 expression levels after 24 hours, indicating that the effect observed after a combined exposure to UV+O₃ was probably due to UV. These data suggest that UV induces inflammatory damage faster than O₃. At DAY 4, even if UV induced a higher increase of COX-2 (+72%), O₃ altered its expression levels (+27% if compared to control tissues) as well. Between-group differences (M89PF treated samples vs pollutant-exposed samples) were all statistically significant ($p\leq0.05$) in favour of M89PF-treated samples (Figure 3B).



Figure 2 Preventive role of M89PF on the increased expression levels of the oxidative marker 4-HNE. 4-HNE expression levels in skin explants samples exposed to O_3 , UV, O_3 + UV and/or pretreated topically with M89PF for 24h and analyzed at DAY I (**A**) or at DAY 4 (**B**). Magnification 40X. Scale bar 100 μ m. Right panels show the quantification of relative immunofluorescence staining for 4-HNE after different treatments (upper panel, DAY I; below panel, DAY4). Data are expressed as the mean of the three different subjects ± standard deviation. *p≤0.05 Ctrl vs Pollutant; [#]p≤0.05 M89PF + Pollutant vs Corre-sponding Pollutant. One-way ANOVA followed by Bonferroni post hoc test.

Moreover, at DAY 1, the expression levels of ASC had significantly increased ($p \le 0.05$) in untreated skin samples after exposure to UV, O₃ and UV+ O₃ (+62%, +102% and +109% respectively if compared to control tissues) (<u>Figure S4A</u>). Pre-treatment with M89PF prevented from pollutant-induced increase in ASC levels. Similarly, between-group differences remained significantly ($p \le 0.05$) in favour of M89PF-treated samples after all exposure exposures and after four days of topical application of M89PF (Figure S4B).

Discussion

Results from this ex vivo study confirm that O_3 and UV alone or combined are deleterious to the upper skin layer.^{6,15,38–40} Exposure to exposome factors significantly (p≤0.05) decrease Filaggrin and Involucrin levels, both markers for the integrity of the natural skin barrier. Moreover, these exposome factors significantly (p≤0.05) increase the levels of oxidative stress markers, such as 4-HNE and AhR, and that of COX-2 and ASC, both being markers of skin inflammation and skin aging.

A single application of M89PF prior to exposure to O_3 and UV alone, or both combined, significantly (p \leq 0.05) reduced the depletion of FLG and INV, thus helping to maintain and protect the natural skin barrier. Moreover, in M89PF-treated samples, the levels of oxidative stress markers triggered by exposure to the exposome factors remained unchanged, compared to negative controls at DAY 1 and DAY 4, with a significant (p \leq 0.05) difference to untreated exposed samples. Similar results were observed for the skin inflammation markers. As mentioned in the Introduction, M89PF contains different active ingredients; in particular, the minerals contained in M89PF reinforce the natural defences of the skin in restoring the natural skin barrier, stimulating antioxidant activity, and reducing



Figure 3 Preventive role of M89PF on the increased expression levels of cyclooxy-genase 2 (COX-2). COX-2 expression levels in skin explants samples exposed to O_3 , UV, O_3 + UV and/or pretreated topically with M89PF for 24h and analyzed at DAY I (**A**) and DAY 4 (**B**). Magnification 40X. Scale bar 100 µm. Right panels show the quantification of relative immunofluorescence staining for COX-2 after different treatments (upper panel, DAY I; below panel, DAY 4). Data are expressed as the mean of the three different subjects ± standard deviation. *p≤0.05 Ctrl vs Pollutant; p=0.05 M89PF + Pollutant vs Corre-sponding Pollutant. One-way ANOVA followed by Bonferroni post hoc test.

inflammation.^{41–46} M89PF has been enriched with probiotic fractions, shown to enhance re-epithelization of the skin and help to limit dysbiosis, inflammation and to restore the natural skin barrier.^{47–49} M89PF has been developed to repair the skin barrier and to reinforce skin defences against exposome factors.

We agree that the present study was conducted ex-vivo and that the sample size may be considered as small. However, results demonstrate that at tissue level, the tested exposome factors have an impact on the upper layers of the skin, and that M89PF provides an unambiguous benefit in protecting these tissues from external aggression. Further research is necessary to confirm the obtained assay results and, potentially, in other skin conditions that may be triggered by exposome, in order to clinically confirm the present observations.

Conclusion

The human body is permanently exposed to O_3 and UV radiations, and further research is necessary to assess the impact of the combined exposure and cumulative effects of different air pollutants that affect each other's response.

M89PF helps to prevent skin barrier damage, as well as oxidative stress and inflammation induced by exposome factors such as UV and O_3 .

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Disclosure

Delphine Kerob is an employee of Cosmetic Active International, a L'Oréal company. The other authors have no conflict of interest to disclose in this work.

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