Sunscreen with the New MCE Filter Cover the Whole UV Spectrum: Improved UVA1 Photoprotection In Vitro and in a Randomized Controlled Trial

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Background: UVA1 rays (340–400 nm) contribute to carcinogenesis, immunosuppression, hyperpigmentation, and aging. Current sunscreen formulas lack sufficient absorption in the 370–400 nm wavelengths range. Recently, a new UVA1 filter, Methoxypropylamino Cyclohexenylidene Ethoxyethylcyanoacetate (MCE) exhibiting a peak of absorption at 385 nm, was approved by the Scientific Committee on Consumer Safety for use in sunscreen products. These studies evaluated, in a three-dimensional skin model and in vivo, the protection afforded by state-of-the-art sunscreen formulations enriched with MCE. Trial design: This study is a monocentric, double-blinded, randomized, and comparative trial. This study is registered at ClinicalTrials.gov with the identification number NCT04865094. Methods: The efficacy of sunscreens with MCE was compared with that of reference formulas. In a three-dimensional skin model, histology, protein, and gene expression were analyzed. In the clinical trial, pigmentation was analyzed in 19 volunteers using colorimetric measurements and visual scoring. Results: MCE addition in reference formulas enlarged the profile of absorption up to 400 nm; reduced UVA1-induced dermal and epidermal alterations at cellular, biochemical, and molecular levels; and decreased UVA1-induced pigmentation. Conclusions: Addition of MCE absorber in sunscreen formulations leads to full coverage of UV spectrum and improved UVA1 photoprotection. The data support benefits in the long term on sun-induced consequences, especially those related to public health care issues.

INTRODUCTION

Exposure to solar UVR is the major environmental aggressor for the skin, responsible for skin cancer. Squamous cell carcinoma has been associated with chronic cumulative exposure all life long, basal cell carcinoma has been associated with both chronic and intermittent intense UV exposure, especially early in life (Milon et al., 2014; Verkouteren et al., 2017), whereas melanoma is associated with intermittent recreational exposure or sunburn history during childhood (Dennis et al., 2008; Gandini et al., 2011).Besides skin cancers, acute or repeated sun exposure contributes to numerous harmful effects such as photoimmunosuppression, viral reactivation, premature skin aging, and pigmentary disorders (Gibbs and Norval, 2013; Ortonne, 1990; Rittie and Fisher, 2015).

Solar UV reaching the earth includes UVB (290–320 nm), UVA2 (320–340 nm), and UVA1 rays (340–400 nm). UVB rays represent 5% maximum of solar UV but induce potentially mutagenic DNA photoproducts and are responsible for erythema, skin pigmentation, photoimmunosuppression, and skin cancers. In turn, UVA1 rays account for >80% of solar UV received on the earth. Less energetic than UVB rays but with higher penetration properties, they can reach the deep dermis. In the last decade, their contribution in photodaging (Wang et al., 2014), photoimmunosuppression (Damian et al., 2011; Matthews et al., 2010), polymorphic light eruptions (Badri and Schlessinger, 2020; Lembo and Raimondo, 2018), herpes simplex virus reactivation (Kerr et al., 2012; York and Jacob, 2010), photocarcinogenesis (de Laat et al., 1997; Ley and Fournier, 2000), and hyperpigmentation (Marionnet et al., 2017; Ravnbak and Wulf, 2007) was pointed out. At the biological level, they lead to ROS generation, induce DNA lesions and mutagenesis in epidermal basal cells, damage skin dermis, and alter the expression of genes involved in a large panel of essential functions and pathways, including cancer, proliferation, apoptosis, development, innate immunity, extracellular matrix, response to stress/oxidative stress, and metabolism.

Abbreviations: FC, fold change; ITA, individual typology angle; KC, keratinocyte; MCE, Methoxypropylamino Cyclohexenylidene Ethox-yethylcyanoacetate; MMP, matrix metalloproteinase; SPF, sun-protection factor

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To limit harmful skin damage due to sun exposure, World Health Organization, United States Environmental Protection Agency, and dermatological academies recommend photoprotective behaviors, including the use of sunscreen with effective UVB and UVA absorption (https://www.who.int/uv/sun_protection/en/; https://www.epa.gov/sunsafety/action-steps-sun-safety; https://www.aad.org/public/everyday-care/sun-protection/sunscreen/how-to-select-sunscreen, respectively). To this end, sunscreen products are composed of UV absorbers whose combination determines the overall absorption profile of the final formula (Daly et al., 2016; Moyal, 2012). Today’s state-of-the-art sunscreens can efficiently filter UVB, UVA2, and UVA1 up to ~370 nm. However, they lack a 370-400 nm sufficient absorption in the UVA1 domain owing to the absence of efficient authorized UV filters in this wavelengths range. To show the benefit of providing the appropriate photoprotection in this UVA1 wavelength range, a proof-of-concept study has been performed using laboratory tools UVA1-absorbing molecules in prototype formulations (Marionnet et al., 2018). On the basis of these mandatory positive efficacy results, the whole process of developing a new UVA1 filter was engaged, taking into account the regulatory (human and environmental safety) and industrial feasibility constraints for molecule selection. Therefore, to our knowledge, a previously unreported cyclic merocyanine UVA1 absorber, Methoxypropylamino Cyclohexenylidene Ethoxyethylcyanoacetate (MCE), was synthesized and enabled coverage of the 360–400 nm wavelengths range (Winkler et al., 2014). MCE dossier was approved by the Scientific Committee on Consumer Safety at the European Commission on the basis of safety and environmental profile and is now listed in the Annex VI of European Union–authorized UV filters. MCE can be used in sunscreen formulations up to 3% (https://ec.europa.eu/health/sites/health/files/scientific_committees/consumer_safety/docs/sccs_o_227.pdf).

This study aimed at investigating the benefit of the MCE filter addition in state-of-the-art sunscreen formulations in UVA1 photoprotection, at the absorption profile level, at the biological level using a three-dimensional human skin model, and in vivo using pigmentation (Persistent Pigment Darkening) as a clinical endpoint.

RESULTS
The MCE filter
The MCE filter exhibited a peak of absorption at 385 nm with an absorption coefficient of 63,052 (investigated in ethanol at 0.02 mM) (Figure 1). The MCE filter showed 100% intrinsic stability in simplex medium, that is, hydroethanolic solution, and stability robustness with regard to different stress, including heat (99% after 2 hours at 90 ºC), acid (96–99% after 0.1 M hydrogen chloride at room temperature or 60 ºC), and oxidant (98–99% after hydrogen peroxide treatment). MCE also showed 98–99% photostability under exposure to solar simulation even in the presence of more oxygen (Tables 1–3).

The MCE filter was added in state-of-the-art formulations to produce formulations with enlarged absorption in the UVA1...
In vitro study
A three-dimensional human skin model was used to compare the efficacy of three formulas: a state-of-the-art formula (Reference vitro), absorbing UVB and UVA up to 370 nm, and Formulas vitro 0.7% MCE and 1.5% MCE containing the indicated percentage of MCE filter, respectively, enabling UVA1 absorption up to 385 nm and 400 nm, respectively (Figure 2a and b).

Cell and tissue change analysis. The formulations were applied topically onto reconstructed skin and sham exposed or exposed to a single UVA1 dose (40, 60, or 80 J/cm²). Skins were histologically analyzed, and fibroblasts were immunostained 48 hours later. As previously described, the biologically efficient dose of 40 J/cm² UVA1 induced epidermal alterations in upper layers and disappearance of fibroblasts even at 45°C. These results indicate that MCE could be further integrated into cosmetic galenic support containing water.

Gene expression analysis. In reconstructed skin, UVA1 exposure induced the modulation of expression levels of inflammatory cytokines and matrix metalloproteinases (MMPs) (Marionnet et al., 2014). These markers were used to test the protection of the three formulas after 40, 60, or 80 J/cm² of UVA1 exposure (Table 4). Forty Joules per square centimeter UVA1 significantly increased the levels of MMP1 and proinflammatory proteins IL-1RA, IL-6, and GM-CSF, taking into account dermal cell mortality. IL-8 amount was also 2.7-fold increased, although this was not statistically different from that of the control. At that UVA1 dose, Reference vitro, compared with unprotected condition, decreased the amounts of secreted MMP1, IL-6, and IL-8, reducing them to control values, and, to a lesser extent, GM-CSF. Reference vitro exhibited good protection against the production of the latter proteins, Formulas vitro 0.7% MCE and 1.5% MCE showed no significant difference from the Reference vitro. At that UVA1 dose, IL-1RA secretion was poorly protected by the Reference vitro, whereas Formula vitro 1.5% MCE exhibited significantly better protection than the Reference vitro. Significant differences could be evidenced at 60 and 80 J/cm² UVA1 between formulas containing MCE filter and the Reference vitro. At 60 J/cm², Formula vitro 0.7% MCE was able to significantly reduce the amounts of MMP1, IL-1RA, IL-6, and GM-CSF (P < 0.05). At 80 J/cm², this formula still significantly reduced MMP1, IL-6, and also IL-8 amounts, whereas Formula vitro 1.5% MCE enabled the significant decrease of all the tested proteins, compared with the Reference vitro (P < 0.05).

Table 1. Intrinsic Stability Performed in a Hydroethanolic Solution

| Temperature | Time   | % Remaining | Aspect              |
|-------------|--------|-------------|---------------------|
| 4°C         | 1 month| 100         | Pale yellow limpid solution |
| 45°C        | 1 month| 100         | Pale yellow limpid solution |
| 4°C         | 2 months| 100        | Pale yellow limpid solution |
| 45°C        | 2 months| 100        | Pale yellow limpid solution |

Abbreviations: MCE, Methoxypropylamino Cyclohexenylidene Ethoxycyclohexenylidene.

Table 2. Acidic and Oxidizing Stability Performed in a Ternary Mixture

| Stress Factor | Concentration | Time | Temperature | Remaining (%) | Loss (%) |
|---------------|---------------|------|-------------|---------------|----------|
| Acid          | 0.5%          | 0 h  | Room temperature | 99           | 1        |
|               |               | 1 h  | Room temperature | 98           | 2        |
|               |               | 1 h  | 60°C         | 96           | 4        |
| Oxidant       | 0.6%          | 0 h  | Room temperature | 99           | 1        |
| 5 V H2O2      |               | 2 h  | Room temperature | 98           | 2        |
| Temperature   | 0.5%          | 2 h  | 90°C         | 99           | 1        |

Abbreviations: H2O2, hydrogen peroxide; HCl, hydrogen chloride; MCE, Methoxypropylamino Cyclohexenylidene Ethoxycyclohexenylidene. MCE was solubilized in a 0.5 g/100 ml ternary water/ethanol/isopropanol mixture with the final concentration of 50/40/10 (v/v/v). Duplicates were put in contact with an acidic or oxidizing solution at controlled temperature and time. Aliquotting and analytical follow-up were performed on an UPLC/DAD Acquity apparatus (Waters, Milford, MA). The results showed that MCE is stable to mild hydrolysis and to H2O2 oxidizer. This means that MCE could be further integrated into cosmetic galenic support containing water, with adjusted pH lower than neutrality and without any issue regarding the presence of oxygen or oxidizing agent.
The modulation ratio was high (R > 10 or R < 0.1) for 8 genes, moderate (5 < R < 10 or 0.2 < R < 0.1) for 11 genes, and low (2 < R < 5 or 0.5 < R < 0.2) for 8 genes. Pre-application of the Reference vitro formula reduced the number of UVA1-modulated genes with 22 out of 27 genes. In addition, in the Reference vitro–treated samples, the intensities of these modulations were reduced compared with those in the nonprotected samples, with only 4, 4, and 14 genes exhibiting high, moderate, and low levels of modulation, respectively. When the Formula vitro 1.5% MCE was applied, only 11 out of 27 (41%) genes were found modulated after UVA1. Moreover, the intensities of modulation were decreased compared with those for the Reference vitro–treated samples, with three, two, and six genes exhibiting high, moderate, and low modulation levels, respectively (Figure 4a). In KCs, the same pattern of gene expression modulation was observed.
modulation was observed. The Reference vitro enabled the decrease in the number of genes differentially expressed by UVA1 exposure and the intensity of modulation, compared with those in the nonprotected reconstructed skins. The use of Formula vitro 1.5% MCE afforded better protection than the use of Reference vitro on the number and intensity of modulations (Figure 4b). The detailed results for the 27 and 24 genes in fibroblasts and KCs, respectively, are shown in Figures 5 and 6. In fibroblasts, the use of the Reference vitro state-of-the-art formula during UVA1 exposure was sufficient to significantly abrogate the UVA1-induced modulation of five genes: JUN oncogene, APCDD1 apoptosis-related gene, IL1 proinflammatory gene, TXNIP oxidative stress response gene, and DNAJB1 stress-related gene. Similar to the Reference vitro, Formula vitro 1.5% MCE was also able to protect the UVA1-induced modulation of these genes. In addition, Formula vitro 1.5% MCE significantly reduced the gene expression modulation of DDIT3 apoptosis gene, CSF2 inflammation-related gene, HMOX1 oxidative stress response gene, HSPA6 stress-related gene, GDF15 extracellular matrix–related gene, and all the genes of the antiviral response family (OAS1, OAS2, MX1, MX2, IFIT1, toll-like receptor gene TLR3, DDX58, GBP2, GBP5, and SAMD9), compared with the Reference vitro. In KCs, Reference vitro enabled to avoid the UVA1-induced gene expression changes of proinflammatory genes (IL1A, CCL20, and ICAM1), gene expression changes of antiviral genes (toll-like receptor gene TLR3, OAS1, and OAS2), and gene expression changes of NQO1 oxidative stress response gene. Formula vitro 1.5% MCE had the same effect as Reference vitro for these genes but was
Table 4. Evaluation of Sunscreen Formulations on the Levels of Secreted Proteins in Culture Medium of Reconstructed Skins Exposed to UVA1

| Amount of Protein in the Condition 0 J/cm², No Formula | MMP1 18.3 ± 4.6 ng/ml | IL-1RA 23.0 ± 5.6 ng/ml | IL-6 0.7 ± 0.3 µg/ml | GM-CSF 54.4 ± 13.6 ng/ml | IL-8 2.1 ± 1.2 µg/ml |
|--------------------------------------------------------|------------------------|-------------------------|----------------------|--------------------------|----------------------|
| 0 J/cm² No formula                                     | 1 ± 0.2                | 1 ± 0.1                 | 1 ± 0.1              | 1 ± 0.1                  | 1 ± 0.2              |
| 40 J/cm² Ref vitro                                     | 2.7 ± 0.9              | 7.1 ± 2.1               | 2.4 ± 1.0            | 3.3 ± 1.2                | 2.7 ± 1.4            |
| Fla vitro 0.7% MCE                                     | 1.3 ± 0.3              | 5.7 ± 1.2               | 1.4 ± 0.3            | 2.5 ± 1.1                | 1.5 ± 0.8            |
| Fla vitro 1.5% MCE                                     | 1.1 ± 0.3              | 4.0 ± 0.8               | 1.4 ± 0.6            | 1.5 ± 0.5                | 2.3 ± 1.5            |
| 60 J/cm² Ref vitro                                     | 5.6 ± 3.8              | 33.1 ± 13.0             | 4.6 ± 3.3            | 10.9 ± 10.1              | 5.9 ± 4.7            |
| Fla vitro 0.7% MCE                                     | 1.4 ± 0.5              | 7.2 ± 2.2               | 1.3 ± 0.4            | 2.4 ± 1.1                | 1.9 ± 0.6            |
| Fla vitro 1.5% MCE                                     | 1.9 ± 0.7              | 7.4 ± 1.4               | 2.1 ± 1.0            | 2.2 ± 0.5                | 3.1 ± 1.0            |
| 80 J/cm² Ref vitro                                     | 11.1 ± 7.6             | 76.5 ± 27.6             | 4.9 ± 2.2            | 14.2 ± 7.9               | 6.2 ± 2.3            |
| Fla vitro 0.7% MCE                                     | 3.8 ± 3.1              | 23.1 ± 18.4             | 1.9 ± 1.3            | 6.8 ± 6.5                | 3.1 ± 1.7            |
| Fla vitro 1.5% MCE                                     | 2.2 ± 1.0              | 12.6 ± 3.5              | 1.4 ± 0.5            | 3.5 ± 2.3                | 2.2 ± 0.8            |

Abbreviations: CI, confidence interval; Fla, formula; MCE, Methoxypropylamino Cyclohexenylidene Ethoxyethylcyanacetate; MMP1, matrix metalloproteinase 1; Ref, reference.

The amount of secreted proteins was measured 48 hours after UVA1 exposure by ELISA in culture media of reconstructed skins protected or not by a Fla. This was divided by the average number of living fibroblasts. Values of control samples (no UV, no formula) are indicated in the first line and were then adjusted to the 1 value. Values are expressed as means of four independent experiments ± 95% CI. The significant difference between 40 J/cm² and 0 J/cm² in the absence of formula is reported in bold (P < 0.05).

1Significant difference from reference vitro (P < 0.05).
2Significant difference from reference vitro (0.05 < P < 0.1) in Student’s t-test.

Figure 4. Evaluation of sunscreen formulations on overall gene expression modulations in reconstructed skins exposed to UVA1. Six hours after exposure to 40 J/cm² UVA1, the level of 27 and 24 transcripts were measured by qPCR in (a) fibroblasts and (b) keratinocytes of reconstructed skins, respectively, protected or not by a sunscreen formula. Three independent experiments were performed. Transcripts levels of UVA1-exposed samples were compared with those of nonexposed samples using a Student’s t-test. A transcript level was considered modulated if the ratio of transcripts level R in UVA1-exposed sample to transcript level in the nonexposed sample was >2 or <0.5 and if the P value was <0.05. Color code gives the intensity of modulation of transcripts level by UVA1. ECM, extracellular matrix; Fla, formula; MCE, Methoxypropylamino Cyclohexenylidene Ethoxyethylcyanacetate; Ref, reference.
also able to significantly decrease UVA1-induced modulation rates of NR4A1, CDKN1A, ODC1, FOSB, GDF15, IFIT1, MX2, OAS1, SLC7A11, and TXNRD1, compared with Reference vitro. Taken together, these results show that the state-of-the-art Reference vitro and Formula vitro 1.5% MCE decreased or abrogated the gene expression modulation induced by UVA1 for most of the studied genes. Formula vitro 1.5% MCE had significantly better protection than Reference vitro against the gene expression modulation induced by UVA1.

The performance of Reference vitro and Formula 1.5% MCE was tested using graphical representation and statistical analysis of the overall gene expression modulation (Figure 7). The biplot visualization allows for a simultaneous representation of the modulation of gene expression and the conditions UVA1, Reference vitro + UVA1, and Formula vitro 1.5% MCE + UVA1. In fibroblasts (Figure 7a) and KCs (Figure 7b), the UVA1 conditions were gathered with all the UVA1-modulated genes (gray and yellow points), except for HSPA6 in KCs (as detailed in Figure 6). On the first horizontal axis, the Formula vitro 1.5% MCE + UVA1 conditions were located at the opposite of UVA1 conditions, whereas the Reference vitro + UVA1 conditions were located in between. These data show that Reference vitro against the gene expression modulation induced by UVA1.

In vivo study
Then, a human in vivo study was performed and statistically analyzed in 19 volunteers to compare the photoprotection efficiency of a state-of-the-art formula (Reference vivo), absorbing UVB and UVA up to 370 nm, with that of Formula vivo 1.5% MCE containing 1.5% MCE filter, absorbing UVB
and UVA up to 400 nm (Figure 2c and d). Both absorption spectra were practically superimposed from 290 nm to 350 nm. Vehicle, Reference vivo, and Formula vivo 1.5% MCE were applied on the back of volunteers before exposure to 50 J/cm² UVA1. Chromametry measurements and visual assessment of skin pigmentation were realized before and 2 hours and 24 hours after UVA1 exposure, corresponding to Persistent Pigment Darkening (Moyal et al., 2000; Sklar et al., 2013). As expected, 2 hours after UVA1 exposure, skin darkening of the vehicle-treated zone was induced, with a decrease in skin luminance ($D_L$) and skin color value ($D_{C14}$ individual typology angle ($D_{ITA}^c$)) and an increase in skin pigmentation ($D_E$), compared with that of the nonexposed zone. These changes were maintained 24 hours after UVA1 exposure (Figure 8a). The use of the Reference-vivo-Formula significantly reduced these changes, at both time points. Compared with Reference vivo, Formula vivo 1.5% MCE enabled a significant increase in $D_L$ and $D_{ITA}^c$ and a decrease in $D_E$, leading to values closer to those of baseline (Figure 8a). Visual assessment on the same tested zones (Figure 8b–d) showed that 2 hours and 24 hours after UVA1 exposure, the vehicle-treated zones presented pigmentation scores between 6 and 10, with 74% of volunteers having a score of 9 or 10 at 2 hours. On the Reference-vivo-formula zone, no more pigmentation score of 10 was observed, the number of volunteers with a score of 9 was drastically reduced, and 5% of the volunteers presented a score of 4. The use of Formula vivo 1.5% MCE reduced even more the number of volunteers with high pigmentation scores and increased the number of volunteers with scores of 4 and 5 (Figure 8b). A statistical analysis on averaged pigmentation scores showed that the use of Reference vivo significantly decreased the mean pigmentation score compared with the use of vehicle and that Formula vivo 1.5% MCE significantly decreased the mean pigmentation score compared with Reference vivo (Figure 8c), as exemplified on the photograph of one subject 24 hours after UVA1 exposure (Figure 8d).

**DISCUSSION**

The whole UV spectrum (100–400 nm) is classified as carcinogenic to humans by the International Agency for Research on Cancer of the World Health Organization.
A growing scientific matter has emphasized the harmful effects of UVA1, the less energetic solar UV wavelengths but representing the largest amount of UV rays reaching the earth. Thanks to the approval by the Scientific Committee on Consumer Safety, the new UVA1 filter, MCE, can now be used in sunscreen products. Owing to its absorption peak at 385 nm, MCE addition in state-of-the-art reference formulas actually enlarged the profile of absorption up to 400 nm.

Using formulas with identical SPFs and similar profiles of absorption up to 370 nm, the efficacy of protection against UVA1 was assessed with or without an MCE filter in the formula.

First, protection was assessed by UVA1-induced epidermal and dermal alterations in a three-dimensional skin model. As expected, under such UVA1 exposure, with realistic doses corresponding to around 2–4 hours of sun exposure depending on latitude and day of the year, the reference sunscreen showed protection of most of the parameters compared with that of unprotected conditions, in line with its absorption profile covering UVA1 up to 370 nm. In this study, on the basis of histological, biochemical, and molecular data, the improvement of protection using formulations containing MCE was demonstrated in a concentration-dependent manner, 1.5% in the formulation being more effective than 0.7%.

UVA1 wavelengths are highly penetrating rays. The protection in the dermis was proven with better protection against dermal fibroblasts alterations and cytokines and MMPs release as well as fibroblastic gene expression.
modulations, especially those related to oxidative stress, inflammation, and dermal matrix organization. These deep cutaneous impacts have been previously linked to photoaging process and solar elastosis formation (Quan et al., 2009), with a specific role of UVA1 rays (Lavker et al., 1995; Tewari et al., 2014; Wang et al., 2014). These insidious biological alterations have to be taken into account with regard to everyday chronic exposure to UVA1 rays, which are present during most part of the day and less affected by seasons than UVB rays (Jablonski and Chaplin, 2010; Tewari et al., 2013). In that respect, the use of MCE in a daily photoprotection would surely improve, in the long term, the prevention of photoaging signs afforded by sunscreens, as shown in long-term follow-up studies (Hughes et al., 2013; Randhawa et al., 2014). Molecular data revealed a maintenance of genome expression, as evidenced by the principal component analysis. The selected genes were representative of various biological functions in both cutaneous compartments.

The major mode of action of UVA1 rays is the generation of ROS, leading to subsequent oxidative damage to cellular components such as DNA, lipids, and proteins. This impact has been monitored using the modulation of expression of several target genes of the Nrf2 pathway—TXNRD1, NQO1, SLC7A11, or HMOX1—as a sensitive indicator of ROS presence (Gegotek and Skrzydlewska, 2015). Their expression was upregulated by UVA1 exposure, and in both cell types, the formula containing MCE significantly better prevented this induction than the reference formula (Figures 5 and 6).

UVA1 rays are major contributors to photoinmunosuppression (Damian et al., 2011; Matthews et al., 2010). Their impact is also associated with modulated genes related to innate immunity, with the expression of antiviral defense genes particularly downregulated and inflammation-related markers upregulated (Marionnet et al., 2014). Immune response and antiviral defense are key issues for clinical consequences, such as Herpes simplex virus reactivation or polymorphic light eruptions (Kerr et al., 2012; York and Jacobe, 2010). A competent immune response is also mandatory in the prevention of skin carcinogenesis, both innate and adaptive immunity participating in tumor immunosurveillance and antitumor effects (Jakobisiak et al., 2003; Valejo Coelho et al., 2016; Yu et al., 2014). This study showed that MCE addition in sunscreen formulation can significantly improve photoprotection compared with the reference, with most genes related to immunity having their gene expression normalized using MCE.
In addition, other genes important in the fate of KCs, the cells at the origin of carcinomas, have their expressions less perturbed when the sunscreen protection is enlarged by adding MCE: notably, the transcription factors NR4A1, exhibiting growth-promoting, angiogenic, and prosurvival activity in most cancers; DDIT3 (CHOP/GADD153) inducible in KCs by genotoxic stress and growth arrest signals such as UV exposure, whose upregulation reflects endoplasmic reticulum stress (Garmyn et al., 1995; Oyadomari and Mori, 2004); and also JUN and FOSB oncogenes. Moreover, the induction by UVA1 of CDKN1A gene, encoding p21Cip1/Waf1 involved in cell cycle arrest to allow DNA repair after DNA damage (Al Bitar and Gali-Muhtasib, 2019), was completely abolished with the use of the formulation containing 1.5% MCE. This could reflect the protection from UVA1-induced DNA damage afforded by the addition of MCE in sunscreens. Altogether, these results are, to our knowledge, previously unreported preliminary biological evidence suggesting a benefit of MCE-containing sunscreen products in UV-induced cancer prevention.

Evaluation of clinical consequences, especially those requiring long-term exposures, is a difficult issue. In this study, the pigmentation corresponding to the Persistent Pigment Darkening was used as a clinical surrogate for the impact of UVA1 exposure because we previously showed that this pigmentation was associated with biological alterations related to inflammation, response to oxidative stress, perturbation of extracellular matrix, cancer, and development (Marionnet et al., 2017). These effects were achieved after exposure to 50 J/cm² UVA1, the same dose as in this study. Such UVA1 dose can be received in <3 hours in summer. In this study, topical application of sunscreens formulations led to a decrease in the level of UVA1-induced pigmentation, with higher prevention of pigmentation when the sunscreen contained 1.5% MCE. On the basis of absorption profiles of both sunscreen formulations, the gain of efficacy of sunscreen containing MCE is attributed to its filtering properties in the 370–400 nm wavelengths range. Because only short-time effects have been assessed through Persistent Pigment Darkening, complementary studies will be engaged to better assess the delayed tanning at longer time points. Although no melanocytes were comprised in the in vitro skin model, making the analysis of direct actors of melanogenesis not possible, some KCs and fibroblasts factors known to contribute to UV-induced pigmentation such as CSF2/GM-CSF, which activates UVA-induced melanogenesis (Imokawa et al., 1996), and GDF15, which has recently been shown to stimulate melanogenesis through MITF/tyrosinase upregulation (Kim et al., 2020), had their mRNA levels significantly normalized when the MCE formulation was used compared with those of the reference formulation.

For the first time, thanks to MCE filtering properties, a full coverage of the whole UV spectrum up to 400 nm was reached, leading to a higher photoprotection against UVA1-induced biological and clinical impacts. The data strongly support the benefits in the long term on sun-induced consequences, especially those related to public healthcare issues. Additional clinical studies should expand proofs of efficacy.

MATERIALS AND METHODS

UVA1 sources
Solar simulators (Oriel, Stratford, CT) with a 1,600 W (in vivo study) or a 1,000 W (in vitro study) Xenon lamp equipped with a dichroic mirror (Oriel, Les Ulis, France) + WG360 2 mm thick filter (Schott, Clichy, France) delivered the so-called UVA1 spectrum (340–450 nm) (Marionnet et al., 2014) (Figure 9).
MCE UVA1 filter
The MCE UVA1 filter (International Union of Pure and Applied Chemistry name: 2-ethoxyethyl (Z)-2-cyano-2-[3-(3-methoxypropylamino) cyclohex-2-en-1- ylidene]acetate; CAS number 1419401-88-9) belongs to the cyclic merocyanine family (Winkler et al., 2014) and has a 322.41 g/mol molecular weight (Figure 1).

Sunscreen products
State-of-the-art formulas were used as references. Formulas vitro 0.7% MCE and Formula vitro 1.5% MCE, containing 0.7 and 1.5% MCE, respectively, were used in vitro. Formula vivo 1.5% MCE, containing 1.5% MCE, was used in vivo. Figure 2 details the formulas compositions, protection factors, and absorption spectra. Owing to formulations’ technical constraints (filters compatibility and stability), the composition of the SPF15 formulas for in vitro and SPF30 formulations for in vivo clinical trial requested specific adjustments in their compositions.

In vitro study

In vitro reconstructed skin model. Normal primary KCs and fibroblasts were isolated from normal human skin as described (Bernerd and Asselineau, 1997). Human skin was obtained from surgical residues after written informed consent from the donors according to the principles expressed in the Declaration of Helsinki and in article L.1243-4 of the French Public Health Code. Reconstructed skins were composed of a dermal equivalent embedding human living dermal fibroblasts, overlaid by an epidermis reconstructed with human epidermal KCs, as previously described (Bernerd and Asselineau, 1997).

In vitro sunscreen application and UVA1 exposure. A total of 2 mg/cm² of sunscreen formula were topically applied onto reconstructed skin samples. Five minutes later, the reconstructed skins were exposed to a single UVA1 dose. During this exposure, Dulbecco’s phosphate-buffered saline without calcium and magnesium (Gibco BRL, Grand Island, NY) medium was used. After exposure, phosphate-buffered saline was replaced by a fresh medium, and reconstructed skin samples were incubated at 37 °C in 5% carbon dioxide for different time periods, depending on the further performed test.

Morphology and fibroblasts counting. Samples were taken and fixed in neutral formalin for histology 48 hours after UVA1 exposure. Histological examination was performed on paraffin sections stained with hematoxylin, eosin, saffron. Fibroblasts were counted using ImageJ software (National Institutes of Health, Bethesda, MA; http://rsb.info.nih.gov/ij/). For each replicate of one condition, six fields were counted. Then, for one condition, the number of counted fibroblasts per replicate was averaged.

Measurement of protein amount (ELISA). The amount of secreted proteins in the culture medium was measured using ELISA 48 hours after UVA1 exposure. MMP1 amount was assessed using Amersham MMP-1 Human Biotrak assay (GE Healthcare, Chalfont St Giles, United Kingdom). Amounts of IL-1RA, IL-6, IL-8, and GM-CSF were simultaneously assessed using the MILLIPLEX Immunology Multiplex assay HCYTOMAG-60K (Merck, Darmstadt, Germany) according to manufacturer’s instructions and were divided by the average number of fibroblasts seen in paraffin sections, as described earlier, to take into account dermal fibroblast number.

Total RNA extraction. Six hours after UVA1 exposure, reconstructed skin samples were rinsed in Dulbecco’s phosphate-buffered saline without calcium and magnesium (Gibco BRL). As previously described, the epidermis was separated from the dermis, and each skin compartment was disrupted and treated with proteinase K. Then, total RNA was extracted from epidermal and dermal samples (Marionnet et al., 2006). The quality of total RNA was analyzed using a 2100 Bioanalyzer and RNA NanoChips (Agilent Technologies, Santa Clara, CA). The amount of total RNA was measured using a Nanodrop ND-1000 spectrophotometer (Thermo Fisher Scientific, Waltham, MA).

Quantitative reverse transcription-polymerase chain reaction (RT-qPCR). First-strand cDNA synthesis was performed using 1 μg of total RNA and the Advantage RT-for-PCR kit (Clontech, Saint Quentin en Yvelines, France). The LightCycler 480 and the LightCycler-FastStart DNA Master SYBR Green kit (F. Hoffmann-La Roche, Basel, Switzerland) were used for qPCR, as previously described (Marionnet et al., 2006). Normalization of mRNA amounts was done using Genorm application and the following housekeeping genes: GAPDH, B2M, RPL13A, RPS28, and RPS9 (Savili et al., 2003; Vandesompele et al., 2002).

Immunostaining. At 48 hours after exposure, immunostaining of vimentin was performed on air-dried vertical 5 μm cryosections using mouse mAb against human vimentin (1:20, Monosan, Unden, The Netherlands). FITC-conjugated rabbit anti-mouse immunoglobulins (1:80, Dako, Glostrup, Denmark) were used as a second antibody. Nuclear counterstaining using propidium iodide was carried out routinely.

Statistical analysis. Fibroblasts number and mean secreted protein amounts were compared using a Student’s t-test. For gene expression, the level of normalized transcripts was compared between nonexposed and exposed samples in the conditions UVA1, Reference vitro + UVA1, and Formula vitro 1.5% MCE + UVA1 using a Student’s t-test. A transcript level was considered as modulated by UVA1 exposure when the R modulation ratio of UVA1-exposed sample to nonexposed sample was >2 or <0.5 and significantly different (P < 0.05).

For the overall comparisons of conditions UVA1, Reference vitro + UVA1, and Formula vitro 1.5% MCE + UVA1, the values of the fold change (FC) were calculated as follows: for FC > 1, FC = R; for R < 1, FC = 1/R. Then, a biplot displaying the first factorial plan of a principal component analysis on the basis of the FC values was carried out. A nonparametric Jonckheere-Terpstra trend test was performed on the first principal component to test the evolution of the FC, depending on conditions.

In vivo study

The full protocol and clinical study report are available at L’Oreal Research Center (Aulnay-sous-Bois, France).

Inclusion and noninclusion criteria. Healthy European volunteers aged 18–40 years with Fitzpatrick phototypes III–IV and ITA ranging from 10° to 35° were recruited. Volunteers without any history of abnormal response to the sun; with no UV tanning marks, freckles, nevi on the back; and without any marks of recent suntan episode were selected. They were requested not to expose themselves to solar or artificial UV sources during the entire study duration.
Demography. A total of 20 volunteers were included, 10 women and 10 men. A total of 10 volunteers were classified as Fitzpatrick phototype III, and 10 were classified as Fitzpatrick phototype IV. The ITA mean of the panel was 23.5°. The panel mean age was 27.85 years, and the median age was 27.5 years.

Protocol. The study was in accordance with the Helsinki Declaration, approved by the Romanian Health Department on 16 December 2013 with registration number N°77373/16.12.2013, and conducted at the investigational site CIDP Biotechnology SRL (Bucharest, Romania) from 31 January 2014 to 20 February 2014. All enrolled volunteers gave their written informed consent.

The study was monocentric, double-blinded, randomized, and comparative; each subject was their own control. The is registered under the ClinicalTrials.gov identification number NCT04865094.

A total of 2 mg/cm² of formulas (vehicle, Reference vivo, or Formula vivo 1.5% MCE) were applied on three zones of 3 × 3 cm² on the volunteers' back before exposure to 50 J/cm² UVA1. Pigmentation was assessed before and 2 hours and 24 hours after UVA1 exposure.

Assignment and masking. The sample size estimation (20 subjects) was performed by a statistician on the basis of a previous study comparing sunscreen formulas under the same UVA1 exposure condition and with similar colorimetric measurements (Marionnet et al., 2018).

At the Day 1 (baseline), each subject fulfilling all inclusion criteria was assigned a randomization number (provided by L’Oréal Research through an electronic case report form) in the chronological order of inclusion. No number was omitted or skipped. The three zones were delimited on the right or left side of the back of the volunteers and were pretreated with vehicle or the two tested formulas in a randomized manner before exposure to UVA1.

Randomization. Randomization list was generated by a computer process before the study by L’Oréal Research and integrated into the electronic case report form. Randomization numbers were attributed to the volunteers following the distribution indicated by the randomization list. The investigator was supplied with sealed envelopes provided by L’Oréal Research for each subject randomization number with information of the formulas (vehicle, Reference vivo, or Formula vivo 1.5% MCE) assignment to each investigational zone. Only in case of safety concerns (local intolerance, adverse event, serious adverse event) was the investigator authorized to open the corresponding envelope. In this case, the volunteer was in major deviation and was excluded from analysis.

Participant flow and follow-up. The 20 randomized volunteers were followed up for 2 and 24 hours after UVA1 exposure and finalized the study. One volunteer was excluded from the statistical analysis owing to the product quantity of application that did not follow the ±2.5% according to the international SPF method (ISO 24444:2019). As a consequence, the data analysis was performed on 19 volunteers (Figure 10).

Safety concerns. No local intolerance, serious adverse events, or adverse events were reported during the study.

Analysis. For colorimetric measurements, a chromameter (CR300, Konica Minolta, Tokyo, Japan) was used to determine L*, a*, b*. Skin color was characterized by the ITA: ITA° = ArcTangent ((L°−50)/b°)180/¡ (Del Bino et al., 2013). The primary criterion was the induced pigmentation (ΔE), between the UVA1-exposed zones and their unexposed adjacent zones, calculated as follows:

\[ \Delta E = \sqrt{(\Delta L^2 + \Delta a^2 + \Delta b^2)} \]

For visual scoring of skin pigmentation, visual pigmentation assessment was done under standard daylight illumination on each exposed zone using a pigmentation scale, grading from 0 (absence of pigmentation) to 13 (pronounced brown pigmentation) (Marionnet et al., 2018).

For statistical analysis, chromometric measurements and clinical grading were statistically analyzed in 19 volunteers using linear mixed models for repeated longitudinal data, with treatment, time, and the interaction treatment X time as fixed factors, the baseline as the covariate, and the subject as a random factor. All within- and between-treatment group comparisons were performed using contrasts on the basis of t-tests. Adjustments for multiple comparisons were carried out using the Benjamini–Hochberg procedure. Statistical analyses were carried out using SAS 9.3 software (SAS Institute, Cary, NC). The two-sided significance threshold was set at 5%.

Data availability statement
This paper does not include large-scale databases (next-generation sequencing or microarray). However, all data generated during and/or analyzed during these studies are available from the corresponding author on reasonable request.
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