

BALANCING SUN EXPOSURE AND SKIN LONGEVITY

Evidence-Based Insights | Suncare Philosophy

A White Paper on the Evolution of Photoprotection, Cellular Defence, and Skin Longevity. Author: Dr. Blake Klug, [MBChB (UCT), Bsc (Hons) Human Genetics (UCT), DipPEC (CMSA)]

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WHITE PAPER

For over three decades, Environ has pioneered the integration of vitamin A and powerful antioxidants into daily skincare, ably supported by intelligent sun science that goes beyond SPF alone.

While SPF remains important for UVB defence, it fails to address the pervasive and chronic damage caused by UVA radiation, oxidative stress, and inflammation.

This white paper substantiates Environ's holistic approach to photoprotection, integrating broad spectrum coverage with antioxidant support and intelligent skincare strategies to preserve skin health and longevity.



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Introduction

Humans have been aware of the deleterious effects of sunlight since the early 1800's which was later then attributed to UV radiation in 1889 [2]. The concept of a sunscreen was first introduced in 1891 by Dr Friedrich Hammer by which he noted that substances that prevent UVR from reaching the skin protect it from sunburn [2]. Since then, there has been a wealth of research which has culminated into the formulations we now consider as modern sunscreens. When a consumer is selecting a sunscreen product they are confronted with a decision, namely selecting the sun protection factor (SPF). Throughout the history of modern sunscreen protection, the emphasis has been on SPF. While SPF has been a valuable benchmark in public sun safety education, its prominence has inadvertently narrowed the public's understanding of what constitutes comprehensive photoprotection in the setting of holistic skincare. There is emerging evidence that a sunscreen product can offer a lot more both in terms of sun protection and skin longevity than SPF alone. As our understanding of skin biology deepens, so too must our approach to sun protection. Modern skincare is shifting away from reactive, symptom-focused products toward preventative, longevity-based solutions. It is no longer enough to prevent sunburn, we must now defend against the silent, cumulative damage caused by daily exposure to environmental stressors that disrupt cellular function, deplete antioxidants, and accelerate the skin's ageing process. By reframing the conversation around sunscreen efficacy—from a focus on SPF values to a more holistic focus on skin longevity, cellular defense, and comprehensive photoprotection—this paper seeks to challenge entrenched perceptions and provide evidence for a paradigm shift in sunscreen development and selection.

UV Radiation and the Skin

For one to understand holistic photoprotection, the stressors on our skin from solar radiation must be examined. Only a portion of the electromagnetic radiation produced by the sun reaches the earth of which the majority is infrared radiation (56%). Approximately 5% is UVA and 0.5% UVB radiation and to lesser degree visible light [3,4]. This represents the energy reaching earth but because UVB photons are far higher energy than UVA photons, the ratio of photons of UVB to UVA is 1:800-1000. That means that every UVB photon that gets blocked is about 10 times greater energy than any UVA photon. With this understanding it becomes obvious that the total energy of UVA reaching our skin is about the same as the energy of UVB. Solar UV radiation (UVR) is classified according to wavelength and is composed of UVA (315-400 nm), UVB (280-315 nm) and UVC (100-280 nm). Only

UVA and UVB reach the earth's surface and therefore human skin since UVC is completely filtered by the Ozone layer. UV penetrates the skin in a wave dependent manner (Fig 1.). About 5-10% of the UVR is high energy shorter wavelength UVB that penetrates the human epidermis and only a small fraction penetrates the upper part of the dermis [3]. UVB radiation has long been recognised as producing immediate visible effects on the skin. UVB exposure induces an inflammatory response in the skin via induction of inflammatory cytokines and vasoactive mediators culminating into what is recognised as 'sunburn' [5,6]. Furthermore, when the exposure to UVB surpasses a critical threshold for inducing damage, keratinocytes initiate apoptotic pathways that ultimately result in cellular demise. These apoptotic keratinocytes can be distinguished by the presence of pyknotic nuclei and are colloquially referred to as 'sunburn cells' [5,7]. UVB additionally contributes to an augmentation of epidermal thickness, a condition known as hyperkeratosis. Through the induction of cellular injury, ultraviolet radiation activates damage response pathways in keratinocytes. Damage-associated signals, significantly modify the physiological state of keratinocytes, facilitating cell cycle arrest, promoting DNA repair mechanisms, and triggering apoptosis in instances of substantial damage. In addition, complex hormonal interactions between keratinocytes and melanocytes mediate melanisation or 'tanning' of the skin. There is upregulation of transcription factors within keratinocytes as a response to UVB induced DNA and cellular damage culminating in melanin production by raising levels of tyrosinase and other melanin biosynthetic enzymes [5]. While apoptosis of damaged keratinocytes serves as a protective mechanism, dysregulation of these processes promote carcinogenesis [8]. UVB-induced DNA lesions, particularly cyclobutane pyrimidine dimers (CPDs), and their mutagenic consequences, has been well established as having mutagenic potential and directly linked to skin carcinogenesis. This is well supported by experimental and clinical data [9,10]. However, there remain challenges in fully elucidating the interplay between DNA repair pathways, mutation spectra, and cellular signalling cascades, with some inconsistencies and gaps in understanding the roles of genetic and epigenetic modifiers. **Nonetheless, one can consider the acute effects of UV radiation on the skin such as sunburn and tanning a 'scar' since these visible changes, while protective, represent complex molecular processes leading to DNA and cellular damage and in some cases contribute to carcinogenesis.**

As mentioned previously 95% of the UVR that reaches our skin is lower energy longer wavelength UVA. Figure 1 demonstrates the wave dependent nature of UVR skin penetration. Longer wavelength UVA can exert its biological effects from the epidermis to the dermis. However, the lower energy properties of UVA radiation have led to the underestimation of its biological effect on the skin [11]. In general, solar radiation is dependent on geographic and environmental factors including time of year, latitude, hour of the day and season. However, UVA is significantly less affected by these parameters when compared to UVB. Furthermore, UVA ominously reaches our skin indoors by virtue of its ability to penetrate through glass, where UVB is almost completely filtered [12].

This emphasises the far greater temporal and depth of exposure of UVA, on our skin and highlights the large component of UVA in the exposome of the skin. It has been well established that UVA induces oxidative stress in all skin compartments from epidermis to dermis. Reactive oxygen species (ROS) are generated after absorption of UVA photons by cellular chromophores leading to lipid peroxidation, protein changes and generation of DNA photoproducts [11,13]. The induction of ROS depletes cellular

antioxidant mechanisms leading to a state of oxidative stress which contributes to deleterious effects on human skin namely, photo-carcinogenesis, pre-mature ageing, immune suppression and hyperpigmentation. [14-19]. In addition, UVR induces pro inflammatory cytokine release from epidermal keratinocytes and prostaglandin, histamine and leukotriene release from mast cells further contributing to an inflammatory state. These ROS-associated stressors promote a state of inflammaging in the skin. Inflammaging is a low grade asymptomatic, chronic inflammation which occurs during physiological ageing and an important factor in the pathogenesis in multiple age related diseases [20]. The clinical manifestations include epidermal and dermal structure and function deterioration represented by patchy/mottled pigmentation, wrinkling, laxity, sagging and dryness. Thus, daily comprehensive sun protection should entail broad spectrum protection against UVA radiation. Protection should target direct UVA absorption as well as the downstream effects thereof. **By replenishing the skins antioxidant mechanisms and bolstering the skins cellular defense systems, the deleterious effects of chronic UVA exposure can be counteracted thus preserving skin appearance, longevity and prevent inflammaging.**

While the detrimental impacts of ultraviolet radiation (UVR) on human skin are well-documented, it is crucial to recognise that ultraviolet radiation also confers certain health benefits. These advantages encompass the synthesis of vitamin D, which is vital for skeletal health and immunological function, as well as potential protective effects against certain diseases affecting the nervous and cardiovascular systems. Furthermore, sufficient exposure to solar UVB radiation may mitigate the likelihood of specific cancers and autoimmune conditions, as reviewed by Queiros et al. [21]. Considering the balance between these advantageous effects and associated risks is fundamental for the formulation of sun protection products.

Sun Protection Factor: Efficacy and Constraints

The Sun Protection Factor (SPF) has historically served as a critical benchmark for evaluating the efficacy of sunscreens, a concept first introduced in 1962. It is formally defined as the ratio of the least amount of UV energy required to produce a minimal erythema on sunscreen-protected skin to the amount of energy required to produce the same erythema on unprotected skin when applied at an even rate of 2 milligram per square centimeter (mg.cm⁻²) of skin [22]. Since shorter wavelength higher energy UVB is responsible for skin erythema or sunburn, **SPF primarily represents UVB protection and does not meaningfully represent UVA protection.** As discussed above, UVA is equally deleterious and thus sun protection products should include UVA protection thereby providing 'broad spectrum' protection. The determination of SPF in a sunscreen product is established in a laboratory under stringent conditions and significant inter-laboratory variability exists [23]. The conditions under which SPF is evaluated are not met in real world application of sun protection products. Ideal application involves, equal application with frequent re application however this has been shown not to be the case. Application by sunscreen users is typically inadequate, applying far less than 2 mg.cm⁻² and several studies have demonstrated that, in practice, users do not uniformly apply sunscreen leading to inhomogeneous product distribution [24].

Furthermore, there is evidence that re-application of sunscreen every 2 hours does not occur among sunscreen users. There are multiple factors that influence SPF product use including weather and anticipated time of exposure [25]. It is plausible that consumers using a higher SPF sunscreen have a false sense of security and may not adhere to good sunscreen application practices leaving them exposed to the adverse effects of UVR. Previously there has been emphasis and promotion of using high SPF sun protection. Notably there is a ceiling of benefit as the numerical SPF designation increases. Specifically, an SPF of 15 absorbs approximately 93.3% of UVB radiation, SPF 30 absorbs 96.7% and SPF 50 absorbs 98% (Fig 2). **It is evident that there is minimal increase in UVB absorption at values beyond SPF 15 [26].** Current Food and Drug Administration (FDA) recommendation is sun-protection product should be SPF 15 or higher and include broad spectrum cover. Interestingly, a study performed on albino mice by Kligman in 1980 determined that a low SPF sunscreen (SPF 2) prevented skin tumourgenesis by 50% where as high SPF sunscreen (SPF 15) completely prevented development of skin cancer [27]. This underscores the fact that SPF 15 was previously regarded as a high level of sun protection, however, manufacturers have subsequently developed formulations with SPF values of 50 or greater. This prompts an inquiry into the rationale behind the pursuit of higher SPF ratings, especially when the incremental benefits appear negligible, and conversely, whether there exist deleterious effects associated with products featuring excessively elevated SPF levels. To comprehend the potential adverse effects, it is imperative to analyse the variables that influence the SPF rating of a product, which will be elaborated upon in the subsequent section. Furthermore, the advantageous effects of solar radiation must also be taken into account, as it is conceivable that excessively high SPF products may attenuate these benefits. Thus, relying on SPF alone as factor when selecting a sun protection product leaves the user at risk of the direct and downstream deleterious effects of UVR.

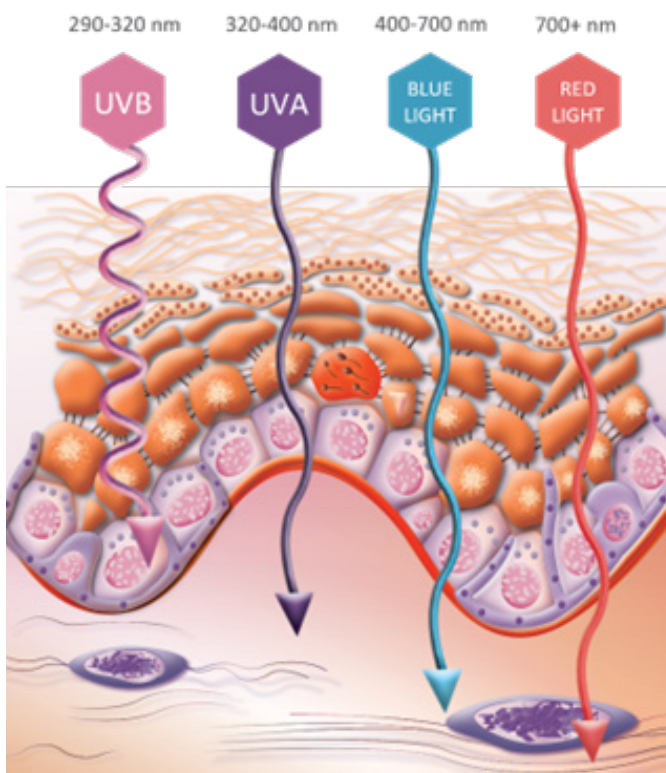


Figure 1. Wave dependent penetration of UVR within the skin.

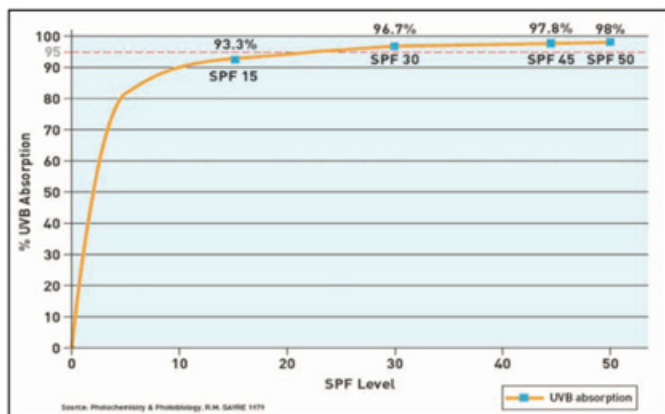


Figure 2. Correlation between UVB Absorption and SPF.

Photoprotection by Ultraviolet Filters

The photoprotection provided by sun protection formulations is primarily achieved by physical barriers that reflect and scatter light and chemical barriers that absorb light, which are referred to as inorganic and organic filters respectively [28] (Fig 3). The mechanism of action for organic sunscreens relies on their chemical composition, which features an aromatic compound conjugated with a carbonyl group. This configuration enables the absorption of high-energy ultraviolet (UV) radiation, resulting in molecular excitation. Upon returning to the ground state, the molecule dissipates this energy by emitting longer-wavelength, lower-energy radiation [28]. Chemical (Organic) sunscreens are typically formulated with agents that are UVA blockers and UVB blockers. As the name suggests, UVB blockers provide protection against higher energy shorter wavelength UVB radiation and act to prevent the deleterious effects described above. Examples include aminobenzoates, cinnamates, salicylates, octocrylene, ensulizole and camphor derivatives. The same is true for UVA blockers, which protect against long-wave ultraviolet A radiation (320–400 nm) and include Benzophenones, Anthranilates, Avobenzone and Ecamsule [28]. Inorganic (physical/mineral) sunscreens include Zinc oxide and Titanium dioxide which act to reflect and scatter light to different degrees based on their reflective index, film thickness, size of the particles and base in which it is dispersed in [28].

UV filters, while crucial for sun protection, have the potential of various adverse effects, ranging from systemic absorption and potential hormonal disruption to dermatological reactions and is well reviewed by Maliyil et al [1] and Jesus et al [29]. Systemic absorption of UV filters Benzophenone-3, Octyl-Methoxycinnamate, and 3-(4-Methyl-Benzylidene) Camphor has been demonstrated. Levels of these compounds have been detected in human serum and urine after a 2 week period of whole body application [30]. Rehfeld et al. investigated the in vitro effects of UV filters on male fertility, hypothesising that these compounds may mimic progesterone. This hypothesis was based on the idea that both progesterone and UV filters can influence calcium (Ca^{2+}) channel signalling, potentially triggering specific biological processes [31, 32]. Further in vitro studies have also investigated the potential endocrine disrupting effects of UV filters in human placental tissue and thyroid activity in an in-vivo rat model [29]. It is important to note that while these agents have not been studied directly in humans, research suggests they may be involved in certain pathological processes, though they are not confirmed to be causative. Nonetheless, this underscores

the need for careful consideration regarding the types and amounts of such substances applied to the skin, particularly given the limited understanding of their long-term effects. UV filters can cause a range of skin reactions, including irritant contact dermatitis, allergic contact dermatitis, phototoxic reactions, and photoallergic reactions. Benzophenones, particularly oxybenzone, is a well-known photo-allergen, causing photoallergic and allergic contact dermatitis. Other reported side effects include dermatitis, dry skin, acne, itching, redness, and general skin irritation [1].

Research has demonstrated that sunscreen ingredients are capable of penetrating not only the superficial layers of the skin but also entering systemic circulation [33,34]. In addition, solution-phase studies have shown that some UV filters can indeed promote the formation of ROS. Such findings have raised concerns about the photogeneration of ROS in sunscreen treated skin. A notable study by Hanson et al. investigated the effects of UV filters such as octocrylene, octylmethoxycinnamate, and benzophenone-3 when applied to the skin surface. The authors observed that, following a period of incubation under UVR, these UV filters not only lost their screening capability but also contributed to an increase in ROS generation within the nucleated cells of the epidermis. Using two-photon fluorescence microscopy and fluorescent probes specific for ROS detection, the study was able to visualise and quantify ROS production at various epidermal depths. The authors do, however, suggest that the simple oil-water emulsifier formulation used in their experiment may have allowed more of each UV filter to penetrate the skin compared to a more advanced formulation that could retain filters on the skin surface [35]. Furthermore, they introduce an idea that inclusion of antioxidants into sun protection formulations could affect UV filter-sensitised ROS generation.

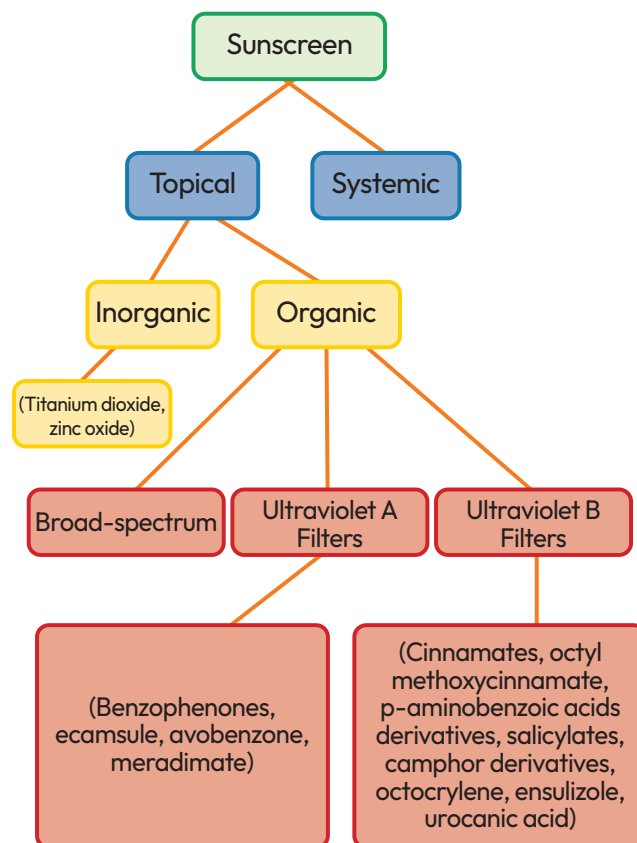


Figure 3. Sunscreen Filters (Maliyil et al).

A large determinant in the formulation strategy to achieve a given SPF is the UV filter type and concentration thereof. Other determinants include, presence of photo-stabilisers, vehicle type and concentration of emulsifiers and particle size of UV filters [36]. Maximum concentrations of a single UV filter that can contribute to a formulation are regulated by FDA, resulting in the use of multiple UV filters especially at higher SPFs [36]. While UV filters offer essential protection against UV radiation, it appears that their use may be not be without consequence, particularly in higher SPF sunscreens. Some UV filters can be absorbed systemically [30, 33, 34], leading to concerns about endocrine disruption[31], carcinogenic potential, various dermatological issues, and cellular damage. Therefore, the formulation of a sunscreen involves a careful balance of risk and benefit—maximising UV protection while minimising potential systemic and dermatological effects associated with increased concentrations and combinations of UV filters.



The Role of Antioxidants in Effective Sun Protection

The downstream deleterious effects of UVR include generation of reactive oxygen species and depletion of the skins natural cellular defence mechanisms thereby promoting a state of oxidative stress and inflammation. Oxidative stress in the skin leads to cellular and DNA damage and triggers the activation of matrix metalloproteinases (MMPs). There is subsequent degradation of structural proteins like collagen and elastin, causing a decline in skin hydration and elasticity, and acceleration of wrinkle formation [37]. Comprehensive holistic sun protection should employ strategies to prevent or minimise UV-induced oxidative damage, boost photoprotection effectiveness, and mitigate skin inflammation.

Defence Mechanisms of the Skin against Oxidative Stress			
Repair Systems	Prevention Mechanisms	Physical Defences	Anti-oxidant Defences
DNA Repair Enzymes	Prevention of ROS production (Chelating agents)	Stabilisation of biological sites (membranes)	Enzymes: - SOD, Glutathione Peroxidase, Catalase Non-enzymes (Low Molecular Weight Anti-oxidants): - Synthesised by the cell - Waste products - Dietary sources (e.g. tocopherols, ascorbic acid, carotene, coenzyme Q10, polyphenols)

Table 1. Skin Defence Mechanisms, modified from Stojiljković et al[40].

Antioxidants are classified according to their mechanism of action and are either primary, secondary or multifunctional [38, 39]. Primary antioxidants, by virtue of their several hydroxyl groups, act directly by donating hydrogen or electrons allowing conversion of free radicals into stable products [39]. There are variety of mechanisms in which secondary antioxidants act and include the chelation of transition metals, singlet oxygen quenching, and restoration of the antioxidant activity of primary antioxidants [39]. Interestingly, some antioxidants are multifunctional and have properties of both primary and secondary antioxidants. The intrinsic antioxidant defence

systems of the skin is robust and include both enzymatic and non-enzymatic mechanisms. Enzymatic systems include the glutathione-peroxidase-reductase enzyme system and superoxide dismutase (SOD) where as non-enzymatic mechanisms involve vitamin E, Vitamin C, glutathione and co-enzyme Q10 [40]. Table 1 highlights the multi-tier complex oxidative stress defence system of our skin and a valuable framework for understanding the skins response to chronic UV exposure.

A strategy to employ is to support the skins natural antioxidant defences that may be overwhelmed by UV radiation. An important metric to consider is the Radical Sun Protection Factor (RSF)—defined as the ratio of the number of free radicals produced in unprotected skin to that of protected skin. It employs an ex-vivo technique using electron spin resonance (ESR) spectroscopy [41,42]. Moreover, it is a biophysical end-point for the protection against ROS induced by UVA and UVB radiation by representing the increase in time required to stay in the sun to generate the same number of free radicals in unprotected skin [43]. The Antioxidant power (AP) of a cosmetic product can also be measured, it is usually an ESR spectroscopy assay that indicates a mixtures capacity to remove a given number of free radicals in certain time intervals and is standardised to vitamin C (Ascorbic Acid). Thus a high AP formulation has the capacity to remove a large number of free radicals in a short period of time [41]. Sunscreens containing vitamin E and vitamin C have been shown to offer significant protection against free radicals from UV exposure [41]. Interestingly, this study noted that there is a significant correlations between UVA protection factor and RSF of a sunscreen product.

Vitamin E is an essential, fat soluble, nutrient that has gained increasing attention in the skincare industry due to its potent antioxidant properties. Unlike certain antioxidants, such as glutathione or ubiquinol-10, which can be synthesised endogenously, cutaneous vitamin E levels depend on dietary intake or topical application. Major natural sources of vitamin E include fresh vegetables, vegetable oils, cereals, and nuts [44]. Among the various forms of vitamin E, x-tocopherol is the predominant homologue found in both mouse and human skin, while x-tocopherol is also present in the epidermis, dermis, and stratum corneum [44]. Interestingly, exposure of human skin to solar-simulated ultraviolet light, comprising of both UVA and UVB wavelengths at sub erythral doses has been show to produce a rapid and substantial depletion of x-tocopherol in human stratum corneum [45]. The authors of this study suggest this depletion is an early and sensitive indicator of photooxidative skin damage. Two pathways are likely responsible for this depletion—direct absorption of UVB radiation and indirect oxidation via singlet oxygen or other ROS generated by UVA. Moreover, the aforementioned processes may also contribute to exhaustion of other endogenous antioxidant systems.

Vitamin E has been identified as the major antioxidant within the skins physiological barrier, the stratum corneum [46]. Compared to the other epidermal layers, the stratum corneum and dermis exhibit limited levels of co-antioxidants such as vitamin C [46] and is likely due to the differences in solubility. Consequently, these regions are particularly vulnerable to oxidative stress, showing the highest levels of protein oxidation following solar UV exposure [47]. Thus, supplementation with vitamin E, alone or in combination with synergistic antioxidants like vitamin C, may enhance the skin’s photoprotective defence and improve

sunscreen efficacy. Extensive evidence supports the role of topical vitamin E in photoprotection and is well reviewed by Thiele et al [44]. Numerous studies have demonstrated that pre-exposure application of vitamin E significantly reduces acute UV-induced skin responses such as erythema, oedema, sunburn cell formation, lipid peroxidation and DNA adduct formation.

Chronic UV exposure outcomes—such as skin wrinkling and tumour formation have likewise been shown to diminish with topical vitamin E treatment. While some debate persists regarding penetration of vitamin E into deeper dermal layers, there is evidence indicating that topical vitamin E can reach these regions. An in vivo study on hairless mice demonstrated that topical administration of α-tocopherol increased vitamin E levels both in the epidermis (62-fold) and the dermis (22-fold) in addition to an increase in enzymatic antioxidant levels [48].

This study, along with others [49], and the well-established photoprotective and antioxidant properties of vitamin E, suggest that it may penetrate into the dermis, where oxidative protein damage predominantly occurs, thereby combatting UV induced inflammation.

Vitamin C (L-ascorbic acid), a water soluble vitamin, is the most prevalent antioxidant found in human skin [50]. Unlike plants and certain animals, humans cannot produce vitamin C because they lack the enzyme L-glucono-γ-lactone oxidase [50]. Even when taken in large oral doses, only a limited amount of vitamin C becomes biologically active and available to the skin [51,52]. As a result, the skin depends entirely on external sources and thus direct topical application could act as an essential source for the skin. **Vitamin C functions as a potent neutralising agent for free radicals by donating electrons within the aqueous compartments of the skin and additionally contributes to the regeneration of vitamin E.** Beyond its role as an antioxidant, it serves as a vital cofactor for essential enzymes involved in collagen biosynthesis and possesses the capability to inhibit the biosynthesis of elastin, thereby mitigating the accumulation of elastin. Furthermore, it attenuates pigment darkening through the inhibition of tyrosinase and sustains skin hydration by safeguarding the epidermal barrier [50,53]. Studies on topical administration of L-ascorbic acid have shown promising photoprotective effects. Early studies by Darr et al on porcine skin revealed protective effects of topical vitamin C against UVB radiation represented by a reduction of sunburn cells [54]. A double-blind placebo controlled study in human subjects reported that topical vitamin C showed a significant reduction in photoaged scores and improvement in wrinkling after application over a 12 week period [55]. This study showed a significant increase in skin hydration and collagen in treated areas compared to untreated skin among subjects. A study by Matsui et al. examined non-erythema endpoints of UV damage and found that adding antioxidants, including vitamin C, to a commercial sunscreen was associated with reduced depletion of Langerhans cells following UV radiation [56]. The depletion of epidermal Langerhans cells, the skin's antigen-presenting cells, has been used as a surrogate for acute and chronic UVR induced immune suppression. The protective effects of topical vitamin C against photoageing, photo carcinogenesis and UV induced immunosuppression is well reviewed by Al Naiami et al [51].

Topical vitamin C positions itself as a potential critical adjunct in comprehensive photoprotection. Its multifaceted biochemical roles, as a potent aqueous-phase antioxidant, a cofactor in

collagen biosynthesis, a modulator of melanogenesis, and an important vitamin E co-antioxidant make it an attractive choice for sunscreen formulations.

Future-Proofing the Skin: A New Era of Sunscreens for Cellular Defence and Longevity

The evolution of sun protection science reflects a progressive understanding of the complex and multifactorial nature of ultraviolet (UV) radiation and its effects on human skin. Historically, emphasis on the Sun Protection Factor (SPF) has shaped consumer perception and regulatory frameworks around sunscreen efficacy. While SPF remains a valuable indicator of protection against UVB radiation, it provides an incomplete assessment of true photoprotection, particularly given the ubiquitous and chronic effects of UVA exposure. UVA, which constitutes the majority of solar UV radiation reaching the earth's surface, penetrates deeper into the skin and drives oxidative stress, inflammation, and the molecular pathways underlying carcinogenesis. **This review underscores the need for a paradigm shift in the conceptualisation of sunscreen efficacy, from a singular focus on SPF to an integrated strategy encompassing broad spectrum protection and antioxidant support.** Broad spectrum formulations, incorporating both UVA and UVB filters, are essential to defend against the full range of photobiological damage. However, the limitations and potential adverse effects of some UV filters, such as systemic absorption, hormonal disruption, and ROS generation, highlight the necessity for balanced formulations. Sunscreen formulation strategies must navigate the demands of optimising ultraviolet protection while concurrently mitigating the possible toxicological and dermatological hazards linked to elevated filter concentrations. There is a wealth of evidence that supports the inclusion of antioxidants as a complementary and protective adjunct in sunscreen formulations. **Vitamins E and C play a pivotal role in neutralising reactive oxygen species generated by UVA-induced oxidative stress.** Their incorporation not only enhances photoprotection but also strengthens the skin's intrinsic defence systems, mitigating inflammation and preserving structural and pigmentation integrity. **The inclusion of antioxidants with UV filters may also reduce filter-induced ROS formation, representing a significant advancement in holistic sunscreen design.**

Recognition of the biological duality of UV radiation, its deleterious effects versus its physiological roles in vitamin D synthesis and immune regulation, demands that photoprotection strategies remain nuanced rather than absolute. **Formulations must strive for balance in order to safeguard against excess UV-induced damage while preserving the minimal exposure required for essential biological processes.** The transition toward holistic photoprotection represents a broader paradigm shift in dermatology and cosmetic formulation. It aligns with the movement toward preventative, longevity-focused skincare, where the goal is not only the avoidance of acute damage such as sunburn but also the preservation of long-term skin health, resilience, and youthfulness. Future metrics of sun protection products should focus on the antioxidant efficacy, through metrics like Radical Sun Protection Factor and Antioxidant Power, in order to elucidate the efficacy of antioxidant adjuncts.

Conclusion

In conclusion, true sun protection is multifactorial, requiring an equilibrium between SPF, UVA defence, antioxidant replenishment, and cellular defence. This integrated approach reflects the next frontier in photoprotection science, shifting from mere prevention of UV-induced erythema to comprehensive maintenance of skin integrity, function, and longevity.

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