Botanicals for photoprotection

Angeli E. Torres, Kevin M. Luk, Henry W. Lim

Photomedicine and Photobiology Unit, Department of Dermatology, Henry Ford Health System, Detroit, Michigan 48202, USA.

Correspondence to: Dr. Henry W. Lim, Department of Dermatology, Henry Ford Hospital, 3031 West Grand Boulevard, Detroit, Michigan 48202, USA. E-mail: hlim1@hfhs.org

How to cite this article: Torres AE, Luk KM, Lim HW. Botanicals for photoprotection. Plast Aesthet Res 2020;7:57.

http://dx.doi.org/10.20517/2347-9264.2020.87

Received: 20 Apr 2020  First Decision: 24 Aug 2020  Revised: 13 Sep 2020  Accepted: 29 Sep 2020  Published: 21 Oct 2020

Academic Editor: Salvador Gonzalez  Copy Editor: Cai-Hong Wang  Production Editor: Jing Yu

Abstract

The importance of photoprotection against the deleterious effects of excessive and chronic exposure to sunlight is now well established. Photoprotective measures include behavioral modifications such as seeking shade, wearing photoprotective clothing, wide-brimmed hat and sunglasses, and applying sunscreen to exposed areas. Data on botanical topical and oral preparations have demonstrated photoprotective potential in in vitro, animal, and human studies. This review will focus on botanicals that have been most extensively studied, namely, Polypodium leucotomos extract, green tea, pomegranate, resveratrol, curcumin, and silymarin. These agents have shown promise in mitigating ultraviolet-induced acute changes on the skin, chronic photodamage, and even skin cancer prevention. However, it must be emphasized that current evidence indicates that these agents should be used as adjunctive measures rather than as a replacement of the photoprotective behavioral modifications described above.

Keywords: Botanical photoprotection, Polypodium leucotomos extract, green tea, pomegranate, resveratrol, curcumin, silymarin

INTRODUCTION

Electromagnetic radiation, including infrared, visible, and ultraviolet (UV) radiation (UVR), have both beneficial and harmful effects on the health of human skin. In particular, UVR exposure plays a significant role in the development of sunburns, photoaging, photoimmunosuppression, keratinocyte carcinomas, and cutaneous melanoma. They can also induce and exacerbate photosensitive dermatoses. Following...
exposure to UVR, reactive oxygen species (ROS) are generated in the skin, which result in oxidative stress. This contributes to acute UV-induced erythema (i.e., photoinflammation) and tanning through upregulation of cyclooxygenase-2 (COX-2), which is involved in early inflammation. With chronic UVR exposure, the formation of photoproducts and ROS can lead to DNA damage, while at the same time cause downregulation of tumor suppressor genes. This allows cells to continue replicating while genetic mutations go unrepaird resulting in cancer formation (i.e., photocarcinogenesis). ROS in the skin can also activate proteins which play an important role in photoaging through blockage of collagen gene transcription, inhibition of collagen synthesis, and overexpression of enzymes that break down collagen.

Photoprotection against UV and visible light is one preventative health strategy to reduce the negative effects of electromagnetic radiation. Historically, photoprotection has been achieved through topical routes. Photoprotection strategies include behavioral modifications such as seeking shade while outdoors, wearing photoprotective clothing including hats and sunglasses, as well as applying sunscreen on otherwise exposed skin sites.

Sunscreens are the most widely recognized means of photoprotection by the public; however, they do have several limitations. These include the need for regular reaplication and lack of efficacy due to under-application (i.e., not applying sufficient amounts). More recently, the ecological safety and potential human toxicity of organic sunscreens, such as oxybenzone and octinoxate, have raised concerns among dermatologists and the general public. However, there is limited evidence of direct toxic effects of organic sunscreens in humans and coral reef species. Nevertheless, inorganic-based sunscreens - namely, zinc oxide and titanium oxide-based sunscreens - have been recommended as an alternative for those concerned about the potential health and environmental impact of organic sunscreens. In addition, alternative photoprotective methods have gained increased interest as adjunct protection against UVR and visible light exposure.

Systemic photoprotection has been used in conjunction with topical photoprotection. It may be administered either subcutaneously or orally. Examples include vitamins, minerals, polyphenols, carotenoids, and α-melanocyte stimulating hormone analog, as well as various plant-based agents that have been reported to yield photoprotective and anti-photocarcinogenic properties. These agents act through their antioxidant, anti-inflammatory, and immunomodulatory effects. The focus of this article will be to review evidence-based systemic and topical botanicals as photoprotective agents.

**BOTANICALS WITH PHOTOPROTECTIVE PROPERTIES**

**Polypodium leucotomos extract**

*Polypodium leucotomos* is a tropical fern belonging to the family *Polypodiaceae*. It is native to Central and South America where it has been used traditionally to treat various skin diseases including psoriasis and atopic dermatitis. At this time, *Polypodium leucotomos* extract (PLE) is the most well studied botanical photoprotective agent. It is commercially available worldwide as an over-the-counter oral supplement. While there are many different preparations of PLE, most of the studies reported in peer-reviewed literature have been done with Fernblock (Cantabria Labs, Madrid, Spain). PLE has antioxidant, anti-inflammatory, immunomodulatory, tumor suppressive, and anti-aging properties. These qualities are mainly attributed to the fern’s high polyphenol content, which is obtained from the leaves. Polyphenols are the most abundant class of antioxidants present in plant-based food and beverages. The polyphenols present in PLE are p-coumaric acid, chlorogenic acid, vanillic acid, caffeic acid, and ferulic acid. Of these, the most powerful antioxidants are ferulic and caffeic acids. It must be emphasized that the concentration of these constituents can vary depending on the PLE preparation. Accordingly, the different PLE preparations may vary in term of their photoprotective ability. This was demonstrated in a study by Gonzalez et al., where the photoprotective activity of six different PLE preparations (including Fernblock) were tested in vitro.
Results showed that Fernblock® was by far the most active (> 5x on fibroblasts and > 3x on keratinocytes) compared to the other five PLE preparations, of which two were found to have almost no activity.

PLE augments the body’s natural antioxidant system and minimizes the accumulation of ROS in the skin. It has the unique ability to scavenge superoxide anion in contrast to conventional antioxidants, which can neutralize singlet oxygen only. As an anti-inflammatory compound, PLE has been shown to suppress UV-induced erythema and reduce cutaneous phototoxicity from photochemotherapy. It can also increase the minimal erythema dose (MED), the UVR dose required for immediate pigment darkening, and the minimal phototoxic UVR dose. One randomized controlled trial found that healthy human subjects who took oral PLE supplements 240 mg twice daily for 60 days were 6 times less likely to have a sunburn, 22 times more likely to have increased MED, and 15 times less likely to exhibit visible erythema post-UVR exposure. These effects were quantified in a later study by Kohli et al. where it was noted that the intensity of UVB-induced erythema decreased by an average of 8% post-PLE supplementation. Similarly, one study which did not utilize Fernblock®, noted increased MED among subjects with skin phototype I-III following once daily intake of an oral supplement containing PLE for 12 weeks. The possible mechanism underlying PLE’s anti-inflammatory properties include the inhibition of transcription factors and cytokines that mediate photoinflammation, namely, tumor necrosis factor (TNF), inducible nitric oxide synthase (iNOS), activator protein 1 (AP-1), and nuclear factor kappa B (NF-κB). In addition, PLE can also reduce the expression of COX-2 and prostaglandin E2, both of which are involved in the initial steps of the inflammatory pathway.

In terms of immunomodulation, PLE has been found to preserve epidermal Langerhans cells that are otherwise depleted as a result of UVR. Its anti-tumor effects were exhibited in mice that were given oral PLE daily as evidenced by increased expression of the tumor suppressor protein p53. PLE has likewise been shown to accelerate extracellular matrix turnover and promote renewal of dermal collagen through inhibition of matrix metalloproteinases (MMPs) and upregulation of tissue inhibitor of metalloproteinase, thereby supporting its role in the prevention of photoaging.

PLE can also provide photoprotection from wavelengths beyond UV. In a study by Mohammad et al., patients taking 480 mg of PLE daily demonstrated a substantial decrease in visible light-induced persistent pigment darkening and delayed tanning. Furthermore, an in vitro study by Zamarrón et al. showed that PLE decreased cell death and collagen degradation in human dermal fibroblasts that were exposed to visible light and near infrared radiation. Additional benefits of PLE include suppression of photodermatoses such as polymorphous light eruption and solar urticaria. It can also be used as an adjunctive treatment for vitiligo and actinic keratosis in combination with phototherapy and photodynamic therapy, respectively.

PLE has relatively minor side effects, including mild to moderate gastrointestinal symptoms and pruritus, which have been reported in a small percentage of patients receiving doses ranging from 120 mg to 1,080 mg daily. No changes in physical examination, vital signs, and laboratory tests (complete metabolic panel and clotting studies) were observed from baseline among patients taking 480 mg daily of PLE for 2 months.

Green tea
Produced from the leaves and leaf buds of Camellia sinensis, green tea is one of the most widely consumed beverages in the world. Its many benefits are primarily due to its polyphenols - otherwise known as green tea catechins (GTC) - of which the most abundant (65%) is epigallocatechin-3-gallate. Similar to other phenolic compounds, GTC has antioxidant, anti-inflammatory, immunomodulatory, and chemopreventive properties.

In vitro studies on human keratinocytes have shown that GTCs can inhibit activation of AP-1 and NF-κB, decrease UVB-induced apoptosis, and stimulate the production of interleukin (IL)-12. IL-12 is postulated...
to play a role in photoprotection since photoinflammation, impaired DNA repair, and tumorigenesis are associated with an absence of IL-12. In mice, topical and oral GTC has been found to protect against photoinflammation and photocarcinogenesis through inhibition of the mitogen-activated protein kinase inflammatory pathway and upregulation of genes involved in DNA repair.

In humans, previous studies have reported that topical GTC reduces photodamage by decreasing production of UVR-induced cyclobutane pyrimidine dimers (CPDs) and visible sunburn. However, GTCs have poor skin penetration when topically applied due to their poor lipid solubility. They are also subject to photodegradation. In contrast, orally administered GTC has been shown to have good skin bioavailability as evidenced by their presence in skin biopsy specimens and skin blister fluid. This was noted after GTC supplementation equivalent to 5 cups of green tea daily for 12 weeks. However, there is conflicting evidence in the literature as to the efficacy of oral GTC as a photoprotectant.

One study showed that women who consumed one liter of green tea daily for 3 months were found to have increased skin elasticity, improved skin texture, decreased transepidermal water loss, and increased cutaneous blood flow. In contrast, a study of healthy, light-skinned (skin phototype I-II) adults revealed no significant differences in UVR-induced erythema, dermal leukocytic infiltration, and induction of cyclooxygenase and lipoxygenase inflammatory pathways among those taking daily GTC supplements (1,080 mg plus 100 mg vitamin C) compared to those in the placebo group. Further studies are needed to reconcile these conflicting evidences.

Pomegranate

The extract from *Punica granatum* or pomegranate is rich in phenolic compounds, specifically, anthocyanins, catechins, and tannins. Significant amounts of these compounds are present in different parts of the fruit but are most concentrated in the peel and juice. At present, pomegranate extract is widely available as an over-the-counter oral supplement or topical formulation, and is often incorporated in commercially sold skin care products.

Pomegranate extract has anti-inflammatory properties and a very potent antioxidant activity - even greater than that of green tea or red wine. It confers photoprotection through inhibition of UV-induced production of free radicals, erythema and burning, DNA damage, cell proliferation, and apoptosis. It can also decrease collagen breakdown.

Murine *in vivo* studies have demonstrated that topical application of pomegranate extract can replenish antioxidants (including catalase, peroxidase, and superoxide dismutase), as well as reduce photoinflammation. This was evidenced by decreased skin edema, epidermal thickening, dermal neutrophilic infiltrates, ornithine decarboxylase, and COX-2. Additionally, topical pomegranate extract has been found to prolong the latency and lessen the multiplicity of skin tumors, thereby supporting its benefit against photocarcinogenesis.

Oral administration of pomegranate extract was found to be protective against UVB-induced skin cancer formation in mice through downregulation of COX-2, iNOS, cyclin D2, and MMPs. Women who consumed 1000 mg of pomegranate extract or 8 ounces of pomegranate juice daily for 12 weeks were reported to have increased MED in a randomized controlled, open-label study.

The anti-aging benefits of pomegranate extract were further investigated in a 2017 study by Kang et al. using orally administered pomegranate juice concentrated powder (PCP) in hairless mice. In this 15-week, placebo-controlled trial, PCP was given at 100, 200, or 400 mg/kg once daily, and 1 hour prior to thrice weekly UVB exposure. The investigators reported dose-dependent decreases in wrinkle formation, skin...
edema, expression of pro-inflammatory cytokines, apoptosis, and MMP activity. Moreover, PCP-treated mice showed increases in water, collagen type I, and hyaluronic acid contents in the skin, indicating a skin moisturizing effect.

**Resveratrol (Grape seed, grape peel, and red wine)**

Resveratrol is a stilbenoid compound belonging to the non-flavonoid class of polyphenols. It is present in grape seeds, grape peels, as well as red wine. In terms of photoprotection, resveratrol has demonstrated anti-oxidant, anti-inflammatory, and anti-tumor effects in several *in vitro*, animal, and human studies.

In an *in vitro* study by Zhou *et al.*[^29], pre-treatment of cultured human keratinocytes with different concentrations of resveratrol prior to UVB irradiation resulted in concentration-dependent increase in cell viability and a decreased rate of apoptosis. Following UVB irradiation, samples that were pre-treated with resveratrol had 15%-52% more viable cells and 15%-22% less apoptosis than non-treated samples. Furthermore, resveratrol pre-treated samples demonstrated a 1.4-fold increase in expression of Bcl-2 (anti-apoptotic protein), and decreased expression of Bax and caspase-3 (i.e., pro-apoptotic proteins) by 52% and 45%, respectively.

In mice, topical application of resveratrol prior to UVB radiation exposure was found to inhibit UVB-induced skin edema, inflammation, generation of ROS (e.g., hydrogen peroxide and lipoperoxides), and induction of COX and ornithine carboxylase[^8]. Meanwhile, oral administration demonstrated anti-tumor effect through alteration of tumor growth factor beta and NF-κB, both of which are involved in cell proliferation and tumorigenesis[^17].

In humans, resveratrol has been found to afford partial protection against UVR-induced photodamage. Wu *et al.*[^30] conducted a study on 15 healthy volunteers who were subject to repetitive UVR exposure from a solar simulator at a dose of 1.5 MED. Results showed that skin sites which were treated with topical resveratrol had less UV-induced erythema and sunburn cell formation compared to placebo (vehicle only) or negative control (no treatment).

**Curcumin (Turmeric)**

Curcumin is the active constituent of turmeric (*Curcuma longa*), a rhizomatous plant native to South Asia that is now grown in many tropical and subtropical regions worldwide. Turmeric is commonly used as a spice, coloring agent, and for various indications in Ayurvedic and traditional Chinese medicine[^8,14].

Previous *in vitro* studies utilizing human keratinocytes and epidermoid carcinoma cells (i.e., squamous cell carcinoma cell line) have found that curcumin decreases UVB-induced apoptosis and inflammation through inhibition of the NF-κB and MAPK pathways[^8,14]. In keratinocytes and fibroblasts, curcumin decreases the expression of MMP-1, which may help reduce the appearance of wrinkles in photoaged skin[^8]. Moreover, curcumin has demonstrated ability to decrease squamous cell carcinoma tumor growth in mice[^14].

Curcumin may also confer protection against UVA-mediated photodamage. According to a study by Liu *et al.*[^31] (2018), pre-treatment with curcumin 2 h prior to UVA exposure prevented accumulation of ROS and restores the innate antioxidant function of human fibroblasts *in vitro*. In addition, curcumin was shown to mitigate UVA-induced apoptosis, inflammation, and collagen degradation.

**Silymarin (Milk thistle)**

Silymarin is an isoflavone derived from the seeds of the milk thistle plant (*Silybum marianum*), which is one of the oldest known medicinal herbs used in traditional European medicine. Nowadays, it is
chiefly marketed as an oral hepatic supplement and has been used to treat hepatitis, alcoholic liver diseases, cirrhosis, and toxin-induced hepatotoxicity. The polyphenols present in silymarin, which is more appropriately termed flavonolignans, are silybin, silychristin, silydianin, isosilybin, and 2,3-dehydrosilybin.[32-34]

The antioxidant and anti-inflammatory effects of silymarin are similar to other phenolic compounds in that it downregulates the UVB-induced generation of ROS, expression of inflammatory transcription factors and cytokines (e.g., TNF, IL-1, and iNOS), and activation of inflammatory pathways including COX and lipoxygenase pathways. As an immunomodulatory compound, silymarin reverses UVB-induced immunosuppression through decreased production of IL-10, which has been found to be elevated in the presence of cancers and are thought to be responsible for the cancer's ability to evade the host's immune response. In addition, silymarin imparts protection against UVB photocarcinogenesis through activation of p53, and reduction of pyrimidine photoproduct formation[34].

In terms of protection against UVA, one in vitro study on human dermal fibroblasts revealed that pre-treatment with silymarin one hour prior to UVA exposure decreased ROS production, DNA strand breaks, activation of MMP-1, and the pro-apoptotic protein caspase-3.[35] In another in vitro study, silymarin and its flavonolignans were found to inhibit the UVA-induced activity of collagenase, hyaluronidase, and elastase, which are respectively responsible for skin wrinkling, loss of hydration, and sagging, indicating an anti-photoaging effect[36].

However, a study by Fidrus et al.[36] (2019) showed contradictory results wherein silymarin pre-treatment of human keratinocytes in vitro 30 min prior to UVA exposure enhanced UVA-induced cytotoxicity in a dose

<p>| Table 1. Most extensively studied botanicals for photoprotection |
|-------------------------------------------|----------------|-------------------------------------|-----------------|-----------------|</p>
<table>
<thead>
<tr>
<th><strong>Botanical agent</strong></th>
<th><strong>Spectrum</strong></th>
<th><strong>Mechanism of photoprotection</strong></th>
<th><strong>Routes</strong></th>
<th><strong>Models</strong></th>
</tr>
</thead>
<tbody>
<tr>
<td>Polypodium leucotomos extract</td>
<td>UVB, VL + UVA1, IR-A</td>
<td>Antioxidant: Augments natural antioxidant system, can scavenge superoxide anion</td>
<td>Oral</td>
<td>In vitro, mouse, human</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Anti-inflammatory: Increases MED, dose required for IPD, and minimal phototoxic dose; inhibits proinflammatory transcription factors, mediators, and cytokines; decreases VL + UVA1 induced PPD and DT</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Immunomodulatory: Preserves eLCs</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Anti-tumor: Increases expression of p53 tumor suppressor gene</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Anti-aging: Downregulates MMP; upregulates TIMP; prevents VL and IR-A induced cell death and collagen degradation</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Other: Suppresses photodermatoses</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Green tea</td>
<td>UVB</td>
<td>Increases skin elasticity and blood flow; stimulates IL-12; decreases apoptosis, CPDs, sunburn, and TEWL; inhibits AP-1, NFKB, MAPK</td>
<td>Topical, oral</td>
<td>In vitro, mouse, human</td>
</tr>
<tr>
<td>Pomegranate</td>
<td>UVB</td>
<td>Increases MED, skin moisture, and tumor latency; decreases inflammation and multiplicity of tumors; inhibits ROS, erythema, burning, DNA damage, cell proliferation, apoptosis, and collagen breakdown</td>
<td>Topical, oral</td>
<td>Mouse, human</td>
</tr>
<tr>
<td>Resveratrol</td>
<td>UVB</td>
<td>Prevents ROS accumulation; decreases apoptosis, inflammation, MMP-1 expression, and SCC tumor growth; inhibits NF-κB and MAPK</td>
<td>Topical, oral</td>
<td>In vitro, mouse, human</td>
</tr>
<tr>
<td>(Grape seed, grape peel, and red wine)</td>
<td></td>
<td>Increases cell viability; decreases apoptosis, erythema, and sunburn cell formation; inhibits COX, ornithine carbamylase, TGF beta, and NF-κB</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Curcumin</td>
<td>UVA, UVB</td>
<td>Prevents ROS accumulation; decreases apoptosis, inflammation, MMP-1 expression, and SCC tumor growth; inhibits NF-κB and MAPK</td>
<td>Topical, oral</td>
<td>In vitro, mouse, human</td>
</tr>
<tr>
<td>(Turmeric)</td>
<td></td>
<td>Increases cell viability; decreases apoptosis, erythema, and sunburn cell formation; inhibits COX, ornithine carbamylase, TGF beta, and NF-κB</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Silymarin</td>
<td>UVA, UVB</td>
<td>Activates p53 tumor suppressor gene; decreases MMP-1 activation, inflammation, ROS, DNA damage, apoptosis, IL-10, and CPDs; inhibits collagenase, hyaluronidase, and elastase</td>
<td>NA</td>
<td>In vitro</td>
</tr>
</tbody>
</table>

UVB: ultraviolet B; UVA: ultraviolet A; VL + UVA1: visible light + ultraviolet A1; IR-A: infrared A; MED: minimal erythema dose; IPD: immediate pigment darkening; PPD: persistent pigment darkening; DT: delayed tanning; eLCs: epidermal Langerhans cells; MMP: matrix metalloproteinase; TIMP: tissue inhibitor of metalloproteinase; IL: interleukin; CPD: cyclobutane pyrimidine dimers; TEWL: transepidermal water loss; AP-1: activator protein 1; NFKB: nuclear factor kappa B; MAPK: mitogen-activated protein kinase; ROS: reactive oxygen species; DNA: deoxyribonucleic acid; COX: cyclooxygenase; TGF: tumor growth factor; SCC: squamous cell carcinoma; NA: not applicable
dependent manner (i.e., less viable cells with higher silymarin doses). In addition, silymarin pre-treated keratinocytes produced higher amounts of CPDs following UVA exposure compared to non-pre-treated keratinocytes. The mechanism for this silymarin-induced phototoxicity is still poorly understood.

Table 1 summarizes the mechanism of photoprotection and spectrum coverage of the botanical agents discussed above.

Others

Other botanicals that have been reported to have photoprotective effects, albeit have not been as rigorously studied, are summarized in Table 2.

CONCLUSION

Botanical-based photoprotection is likely to increase in popularity as consumer trends worldwide continue to place an emphasis on naturally occurring compounds used solely or in conjunction with synthetic products. The botanicals reviewed above currently have the most evidence available and can serve as options for providers to recommend to patients. These oral and topical botanical products act through a variety of biologic mechanisms to confer protection against the adverse effects of UVR. However, unlike sunscreens, botanical products are not subject to FDA regulations and so rigorous efficacy and safety testing through large-scale controlled therapeutic trials are lacking for many of these agents. As such, their true photoprotective benefit compared to established measures like seeking shade, donning UV-blocking garments, or organic or inorganic topical sunscreens remains to be verified. In addition, the stability of botanical ingredients as well as the optimal concentration of their constituents is unregulated. Therefore, while evidence on their use as an adjunctive means of photoprotection appears favorable, they should be used in conjunction with, and not as a replacement of, pre-existing photoprotection recommendations. Finally, as the biologic effects of other wavelengths of electromagnetic radiation such as visible and infrared ranges continue to be elucidated, it will be critical for future research to evaluate the potential applicability of botanicals for protection in that realm as well.

DECLARATIONS

Authors’ contributions

Made substantial contributions to literature search and writing of initial manuscript: Torres AE, Luk KM
Contributed to writing and editing of manuscript: Lim HW
Availability of data and materials
Not applicable.

Financial support and sponsorship
None.

Conflicts of interest
Torres AE and Luk KM declared that there are no conflicts of interest; Lim HW is an investigator for Incyte, L’Oréal, Pfizer, PCORI, has served as consultant for Pierre Fabre, ISDIN, Ferndale, and Galderma, and has participated as a speaker in general educational session for Johnson & Johnson, and Ra Medical System.

Ethical approval and consent to participate
Not applicable.

Consent for publication
Not applicable.

Copyright
© The Author(s) 2020.

REFERENCES