

Protective effect of different antioxidant agents in UVB-irradiated keratinocytes

Sara Salucci,¹ Sabrina Burattini,¹
 Francesca Buontempo,²
 Alberto Maria Martelli,²
 Elisabetta Falcieri,¹ Michela Battistelli¹

¹Department of Biomolecular Sciences,
 University of Urbino “Carlo Bo”

²Department of Biomedical and
 Neuromotor Sciences, University of
 Bologna, Italy

Abstract

Skin cells can respond to UVB-induced damage either by tolerating it, or restoring it through antioxidant activation and DNA repair mechanisms or, ultimately, undergoing programmed cell death, when damage is massive. Nutritional factors, in particular, food antioxidants, have attracted much interest because of their potential use in new preventive, protective, and therapeutic strategies for chronic degenerative diseases, including skin inflammation and cancer. Some polyphenols, present in virgin olive oil, well tolerated by organism after oral administration, show a variety of pharmacological and clinical benefits such as antioxidant, anti-cancer, anti-inflammatory, and neuro-protective activities. Here, the protective effects of antioxidant compounds against UV-induced apoptosis have been described in HaCaT cell line. Human keratinocytes were pre-treated with antioxidants before UVB exposure and their effects have been evaluated by means of ultrastructural analyses. After UVB radiation, a known cell death trigger, typical apoptotic features, absent in control condition and in antioxidant alone-treated cells, appear. An evident numerical decrease of ultrastructural apoptotic patterns and TUNEL positive nuclei can be observed when natural antioxidants were supplied before cell death induction. These data have been confirmed by molecular investigation of caspase activity. In conclusion, this paper highlights antioxidant compound ability to prevent apoptotic cell death in human keratinocytes exposed to UVB, suggesting, for these molecules, a potential role in preventing skin damage.

Introduction

The skin is the largest organ of the body, comprises a surface area of approximately 1.5-2.0 m² and protects the underlying tissues through melanogenesis, by acting as an effective barrier against the detrimental effects of environmental, chemical and physical agents.¹

Exposure to solar UV radiation is considered a major source of damage for human skin, it induces several harmful responses, including erythema, immunosuppression, edema, sunburn, keratinocyte hyperplasia, wrinkling, dryness, mottled pigment abnormalities, premature aging and skin cancer.^{2,3}

UV radiation (100-400 nm wavelength) is composed of UVA (320-400 nm), UVB (280-320 nm) and UVC (200-280 nm) rays. UVB radiation represents the major cause of serious skin disorders and genotoxic damage.^{4,5} Exposure of mammalian skin to UV light impairs antioxidant defense mechanisms and increases the cellular levels of reactive oxygen species (ROS), which, in turn, damage lipids, proteins and nucleic acids, thus contributing to the process of photocarcinogenesis and photoaging.^{6,7} In addition, exposure of cells to UVB radiation results in the loss of keratinocyte viability, an increase in membrane blebbing, cytoskeletal molecular changes, and apoptosis.⁸ Apoptosis is a well defined, highly regulated form of programmed cell death that is evident in normal skin (*e.g.*, during keratinocyte differentiation) as well as in pathological conditions. UVB-induced apoptosis is mediated by apoptotic signals, activating two main pathways: intrinsic (mitochondrial) and extrinsic (death receptor-dependent).⁹

This paper discusses the effect of well-known antioxidant molecules against UVB radiation in a keratinocyte cell line (HaCaT). The compounds, chosen for their ROS scavenger and anti-apoptotic properties in keratinocytes, are Melatonin (Mel), Creatine (Cr), Hydroxytyrosol (HyT), Tyrosol (TyR) and, for the first time, Hydroxytyrosyl Laurate (Laur-HyT) and Hydroxytyrosyl Myristate (Myr-HyT).¹⁰⁻¹⁴

Mel, a pineal hormone derived from tryptophan metabolism, is known for its antioxidant and anti-apoptotic action against UVB damage.^{11,14} It was originally identified as the main secretory product of the pineal gland, later demonstrated to be synthesized in a variety of extrapineal sites, with bioactivity vs a number of targets in single cells, animals and humans.¹⁵ Mel can act as a receptor-independent antioxidant,¹⁶ as an antiaging substance and as an anticar-

Correspondence: Michela Battistelli,
 Department of Biomolecular Sciences,
 University of Urbino “Carlo Bo”, Via Ca’ le
 Suore 2, 61029 Urbino (PU), Italy.
 Tel.: +39.0722.304269.
 E-mail: michela.battistelli@uniurb.it

Key words: Keratinocytes; UVB radiation;
 natural antioxidants; ultrastructure; TUNEL.

Acknowledgements: this work was supported by the University of Urbino “Carlo Bo”, in particular by the “Progetto di Valorizzazione DISB 2017”. We would like to thank Dr. Francesco M. Giordano and Dr. Debora Burini for their skillful technical assistance.

Received for publication: 8 March 2017.

Accepted for publication: 28 July 2017.

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European Journal of Histochemistry 2017; 61:2784
 doi:10.4081/ejh.2017.2784

cinogenic factor within a wide range of concentrations.¹⁷

Cr is a molecule with protective effects in neurological and atherosclerotic diseases, in brain and nerve traumatic injury and in UV-induced mitochondrial DNA mutations in skin cells¹⁸ and in skeletal muscle cells after H₂O₂ exposure.¹⁹

HyT, a natural antioxidant polyphenol¹³ present in virgin oil, shows a variety of pharmacological effects, such as antioxidant, anti-cancer, anti-inflammatory and neuro-protective activities.²⁰

HyT, obtained by chemical or enzymatic hydrolysis of oleuropein glucoside, or by chemical synthesis, is the main biophenol in olive oil, regularly assimilated in humans.

TyR, a liposoluble, non-carboxyl mono-phenol, is one of the most representative phenols in olive leaves and in extra-virgin olive oil. Its antioxidant and anti-proliferative activities, as well as its ability to inhibit pro-oxidation processes on human LDL particles,²¹ have been demonstrated.

Because of the limited solubility of HyT and TyR in lipid media, the search for new lipophilic derivatives with enhanced properties is of great interest. Several authors modified the chemical structure of these compounds increasing the alkyl chain and obtaining the ester or ether derivatives. Mateos *et al.* and Madrona *et al.*^{22,23} demonstrated that HyT derivatives with short and medium acyl side chains, such as the acetyl, butyryl and lauryl, are much more effective

in protecting the RBCs against AAPH induced hemolysis than HyT. The greatest efficiency was obtained when the alkyl chain has a length around 12 C-atoms (Laur-HyT). Moreover, Burattini *et al.*¹³ demonstrated that Laur-HyT has a protective antioxidant effect against H₂O₂ treatment, having a role in the prevention of apoptotic death in normal and tumor cells.

For that, in this work, Laur-HyT (12 C-atoms) and Myr-HyT (14 C-atoms) activities against UVB rays, have been analyzed to understand if the antioxidant effect also occurs in the presence of different number and position of the phenolic hydroxyl groups.

Materials and Methods

Cell culture

HaCaT cells, spontaneously immortalized human keratinocytes, were grown in DMEM supplemented with 10% heat-inactivated fetal bovine serum, 2 mM glutamine and 1% antibiotics and were maintained at 37°C in humidified air with 5% CO₂. Cell behavior was progressively monitored by means of inverted microscope.

Apoptosis induction and antioxidant administration

For apoptosis induction cells were cultured to 80% confluence and washed with PBS, covered with a thin layer of PBS and exposed to UVB for 10 min. The irradiating source consisted of a lamp generating UVB light in the range of 290-320 nm with an emission peak at 312 nm (Transilluminator 2000, Bio-Rad Laboratories, Hercules, CA, USA). After the addition of fresh medium, cells were further incubated for 2 h.

Different aliquots of HaCaT cells were

treated for 24 h with 100 μM Mel, or 30 μM Cr monohydrate, or 20 μM HyT, or 5 μM TyR, or 5 μM Laur-HyT, or 5 μM Myr-HyT before UVB exposure. Antioxidant concentrations have been chosen since they did not affect HaCaT cell viability and proliferation as demonstrated in our previous works.^{14,24}

Mel (Sigma-Aldrich, St. Louis, MO, USA) was first dissolved in absolute ethanol at the initial concentration of 100 mM, and then diluted at final 100 μM concentration in culture medium. Cr (Sigma-Aldrich) was dissolved in complete culture medium at final 3 mM concentration. HyT and TyR were recovered with 95% purity from olive oil wastewaters, and further purified by column chromatography. HyT ester derivatives (Laur-HT and Myr-HyT) were prepared according to Mateos *et al.*²² For all treatments, morpho-functional analyses have been performed after 2 h post incubation, which follows UVB radiation.

Scanning electron microscopy (SEM)

Control and treated cells were directly processed on coverslips in Petri dishes.

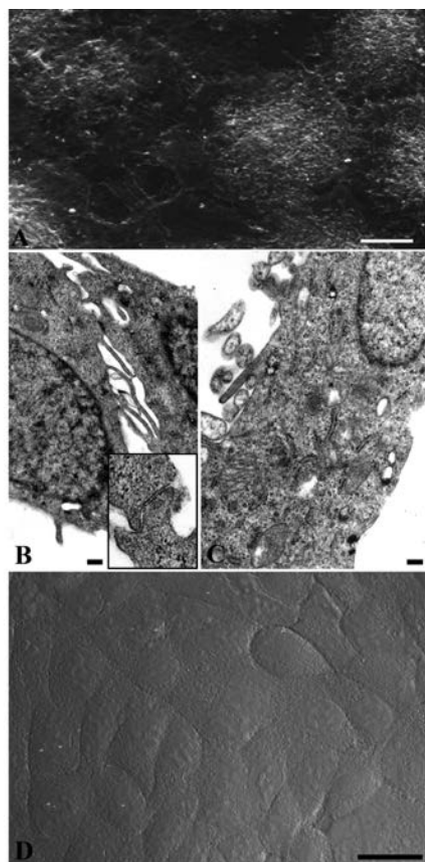


Figure 1. Control cells observed at SEM (A), TEM (B) and CLSM (C). Cells appear mostly polygonal, flattened (A, B) with preserved RER and mitochondria (C) and frequent close cell contacts (B, inset). After TUNEL reaction no labelled nuclei can be observed (D). Scale bars: A) 10 μm; B,C) 0.2 μm; inset in B) 0.1 μm; D) 25 μm.

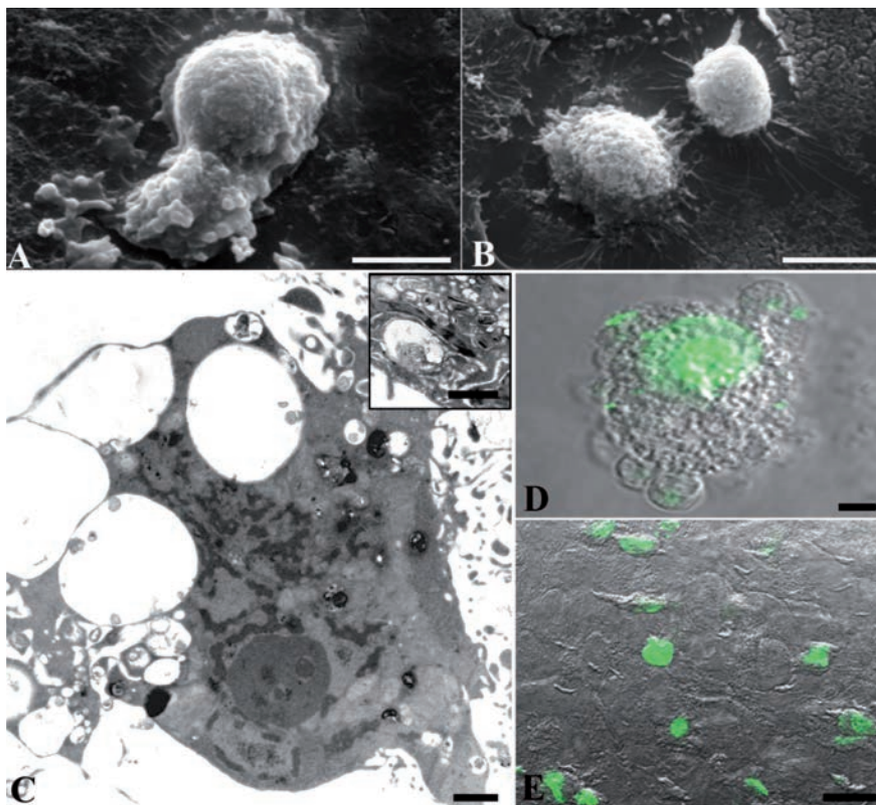


Figure 2. HaCaT cells exposed to UVB and observed at SEM (A,B), TEM (C) and CLSM (D). Apoptotic features, as well as a diffuse positivity to TUNEL reaction, appear. Autophagic vacuoles occasionally appear at TEM observation (C, inset). Scale bars: A,B) 10 μm; C and inset) 1 μm; D) 5 μm; E) 25 μm.

After careful washing, monolayers were fixed with 2.5% glutaraldehyde in 0.1M phosphate buffer for 1 h. All the specimens were post-fixed with 1% OsO₄ in the same buffer for 1 h. After alcohol dehydration, samples were critical point dried, gold sputtered and observed with a scanning electron microscope.²⁵

Transmission electron microscopy (TEM)

HaCaT cells were washed, immediately fixed *in situ* with 2.5% glutaraldehyde in 0.1 M phosphate buffer for 15 min, gently scraped and centrifuged at 1200 rpm. The pellets were fixed in 2.5% glutaraldehyde for additional 30 min.

Cells were post-fixed in 1% OsO₄ for 1 h, alcohol dehydrated and embedded in araldite.^{26,27} Thin sections were stained with uranyl acetate and lead citrate and analyzed at transmission electron microscope.

TUNEL

Control and treated cells, directly processed on coverslips in Petri dishes, were washed and fixed with 4% paraformaldehyde in PBS, pH 7.4, for 30 min, rinsed with PBS and permeabilized with a 2:1 mixture of ethanol and acetic acid for 5 min at -20°C. For the TUNEL technique, all reagents were part of the ApopTag Plus® kit (D.B.A., Dallas, TX, USA) and the procedures were carried out according to the manufacturer's instructions, as described by Salucci *et al.*¹⁴ Specimens were observed with a Leica TCS-SP5 connected to a DMI 6000 CS Inverted Microscope (Leica Microsystems CMS GmbH, Mannheim, Germany); excitation was at 488 nm and emission signals were detected at 517 nm.

Trypan blue (TB) exclusion assay

TB exclusion assay was performed to identify living cells and quantify dead cell number percentage in all experimental conditions. Differences in the percentages of viable cells among groups were determined using one-way analysis of variance (ANOVA), followed by Tukey HSD *post-hoc* tests to evaluate individual group differences. Significance was set at P<0.05. Data were collected from three independent experiments.

Western blotting

Protein assay was performed using the Bio-Rad Protein Assay according to the manufacturer's instructions. Cells were lysed at 10⁷/mL in RIPA lysis buffer (Thermo Scientific, Waltham, MA, USA). Lysates were sonicated to shear DNA and

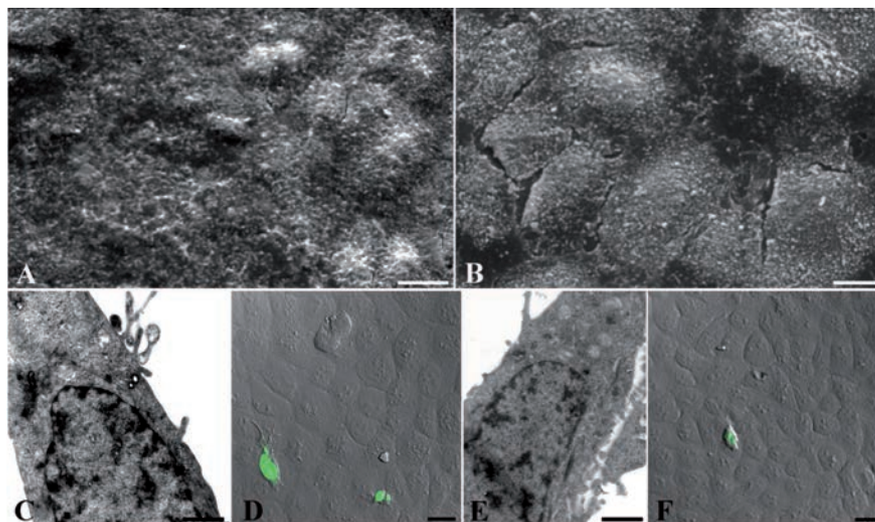


Figure 3. Mel (A,C,D) and Cr (B,E,F) treated cells before UVB exposure and observed at SEM (A,B), TEM (C,E) and CLSM (D,F). All compounds induce an evident apoptotic pattern numerical decrease. Scale bars: A,B) 10 µm; C,E) 0.5 µm; D,F) 25 µm.

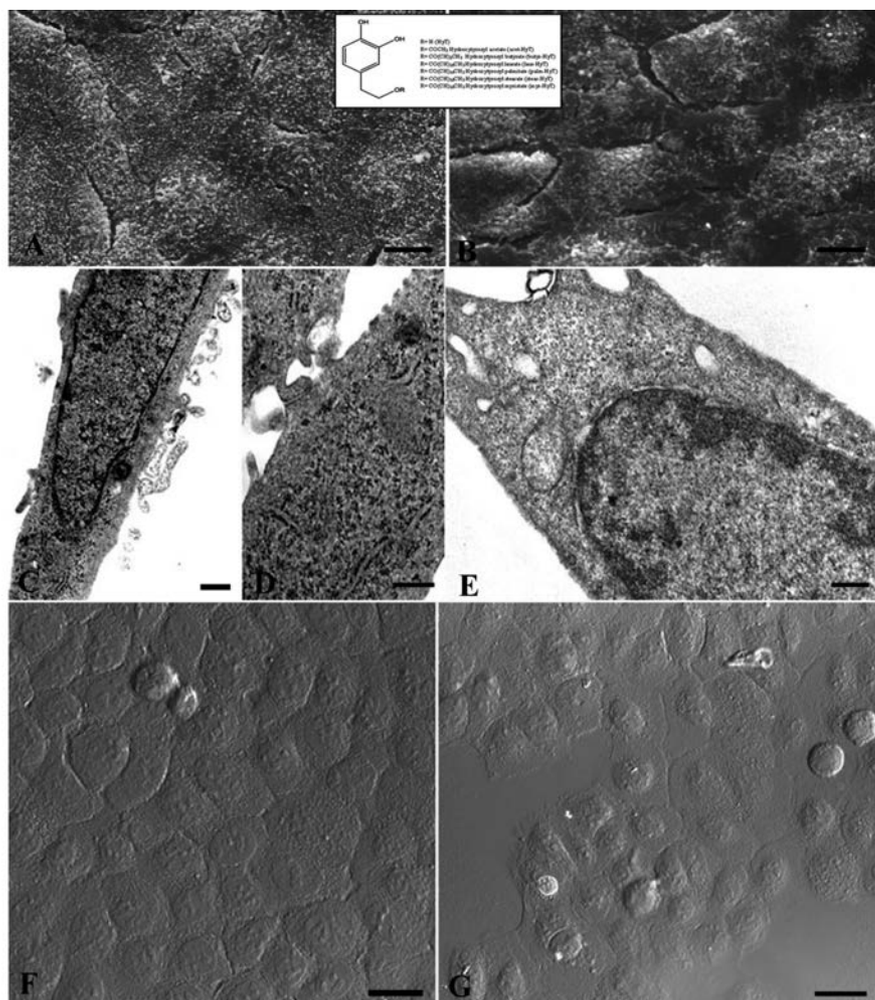


Figure 4. HyT (A,C,D,F), TyR (B,E,G) treated cells before UVB exposure and observed at SEM (A,B), TEM (C,D,E) and CLSM (F,G). Scale bars: A,B) 10 µm; C) 0.5 µm; D,E) 0.25 µm; E,G) 25 µm.

boiled for 5 min with reducing sample buffer. Protein separated (25 µg) on sodium dodecylsulphate polyacrylamide gel electrophoresis (SDS-PAGE) was transferred to nitrocellulose membranes using a semidry blotting apparatus. Membranes were saturated for 60 min at room temperature in blocking buffer, then incubated overnight at 4°C with primary antibody for caspase-8, -9, -3 and PARP (Cell Signaling Technology, Danvers, MA, USA). After washing in PBS containing 0.1% Tween 20, samples were incubated for 1 h at room temperature with peroxidase-conjugated secondary antibody diluted 1:2000 in PBS-Tween 20, and washed as above.²⁴ Bands were visualized by the ECL method and images were achieved through ChemiDoc-it2 Imager (UVP, Upland, CA, USA). Densitometric analysis of cleaved caspase western blotting bands has been performed. Data are expressed as ratio between mean optical density (OD) of cleaved caspase band and mean OD of actin band (protein control), in arbitrary units.

Results

Cells treated with Mel, Cr, HyT, Tyr, Laur-HyT and Myr-HyT alone appeared very similar to control ones, according to quantitative analyses previously reported,^{10-14,17,19} indicating that these compounds did not affect cell viability and/or proliferation. Moreover, TUNEL reaction appeared negative in all conditions (*data not shown*).²⁸

HaCaT cells exposed to UVB radiation

SEM and TEM observations showed how keratinocytes appeared in control condition (Figure 1). Cells were mostly polygonal, flattened and closely confluent (Figure 1A), and displayed a large nucleus with diffuse chromatin (Figure 1B). Close cell contacts (Figure 1B, inset) frequently appeared, as expected for epidermal origin cells. In the cytoplasm well preserved RER and mitochondria (Figure 1 B,C) could be observed. No DNA fragmentation could be revealed in control condition, as confirmed by TUNEL reaction negativity (Figure 1D).

Differently, UVB irradiated samples revealed deep changes in monolayer organization, as well as significant surface and inner cell ultrastructural alterations (Figure 2). Cell monolayer showed a significant decrease in cell confluence, with cell detachment and the appearance of empty spaces, correlated to cell rounding and blebbing (Figure 2 A,B). Apoptotic patterns,

such as chromatin margination and condensation, as well as diffuse cytoplasmic vacuolization, have been detected (Figure 2C). TEM analysis also revealed numerous autophagic vacuoles (Figure 2C, inset). In addition, after UVB radiation exposure, several TUNEL positive, strongly fluorescent nuclei have been detected (Figure 2F). In Figure 2D, a higher magnification of a single cell could be observed where it is possible visualized membrane blebs and DNA cleavage. As also quantified through TB assay, a massive damage appeared after UVB exposure, generating about 60% dead cells if compared to control condition.

Mel and Cr action

Mel (Figure 3 A,C,D) or Cr (Figure 3 B,E,F) treatments, before cell death induction, caused an evident attenuation of apoptotic patterns. A better morphology was observed in all experimental conditions and

TUNEL positive nuclei underwent an evident numerical decrease (Figure 3 D,F). After Mel and Cr pre-treatment cells appeared similar to the control ones. SEM observations showed polygonal, flattened and closely confluent cells (Figure 3 A,B). At TEM, well preserved RER and mitochondria could be evidenced. Nuclear chromatin appeared diffuse both in Mel and in Cr pre-treated cells (Figure 3 C,E). The effects of pre-treatment with Mel and Cr, known in the literature on various cell lines, can be so confirmed in HaCaT cells too and have been considered by Authors a control condition.

Hyt, Tyr and ester derivative action

Also with Hyt (Figure 4 A,C,D,F) or Tyr (Figure 4 B,E,G) pre-treatment, a good preservation of cell morphology could be observed: cells appeared polygonal, flattened and confluent at SEM observation

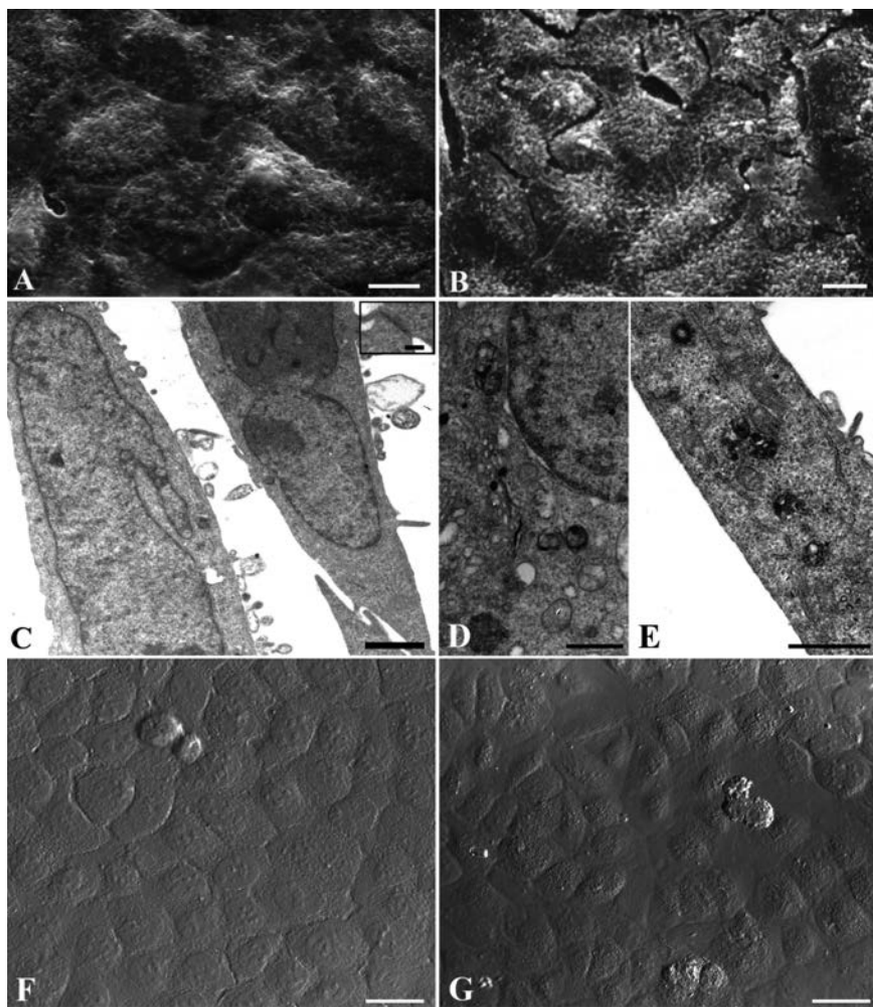


Figure 5. Laur-HyT (A,C,F), Myr-HyT (B,D,E,G) treated cells before UVB exposure and observed at SEM (A,B), TEM (C,D,E) and CLSM (F,G). Scale bars: A,B) 10 µm; C,D,E) 1 µm; F,G) 25 µm; inset of C) 0.1 µm.

(Figure 4 A,B). The cytoplasmic organelles were preserved (Figure 4 C,E) and intracellular junction could be appreciated (Figure 4D). Moreover, few positive TUNEL nuclei appeared after Hyt (Figure 4F) and Tyr (Figure 4G) pre-treatment. Even pre-treatment with Laur-HyT (Figure 5 A,C,F) and Myr-HyT (Figure 5 B,D,E,G) produced good results, at SEM observation we have seen the typical morphology of control cells. At TEM close cell contacts (inset in Figure 5C) could be frequently observed. In the cytoplasm we observed a well preserved RER and mitochondria (Figure 5 C).

TB assay and molecular analyses

Antioxidant protection has been confirmed and quantified by TB assay (Figure 6) evidencing the ability of these compounds to counteract cell death induced by UVB radiation, sensibly decreasing dead cell number. In particular, oil polyphenols appear the better candidates to prevent UVB damage. A well preserved cellular condition in control samples and a cellular

viability decrease after UVB exposure appeared. Interestingly, the antioxidant pre-treatments were able to significantly reduce cell death ($P < 0.01$). In Cr pre-treated samples the dead cell number reduction was significant too, but to a lesser extent ($P < 0.05$). To evaluate anti-apoptotic action of all compounds against UVB radiation, caspase involvement has been evaluated through western blotting analysis. Densitometric analyses showed that caspase-8, -9, -3 and PARP activation, absent in control condition, could be observed after UVB exposure while it was reduced by the different pre-treatments. In Figure 6, the densitometric analyses demonstrated the ability of antioxidants to reduce caspase activation which appeared up-regulated after UVB exposure.

Discussion

This work demonstrates that Mel, Cr, HyT, Tyr, Laur-HyT, and Myr-HyT are able

to prevent UVB-induced apoptosis in a keratinocyte cell line. A massive DNA damage can be inhibited by antioxidant administration, able to significantly reduce cell death, to prevent ultrastructural apoptotic patterns appearance as well as DNA fragmentation, and to reduce caspase activation. All scavengers used in this work, with particular attention to Mel, HyT and Tyr which showed the major anti-apoptotic properties, could represent potential tools in the treatment of UVB exposure-induced skin disorders. As demonstrated in this work, TUNEL reaction evidenced in human keratinocytes after UVB radiation the presence of *in situ* DNA fragmentation, a typical apoptotic feature. All pre-treatments are able to reduce DNA cleavage and, as consequence, to prevent apoptotic cell death.

In particular, we demonstrate that Mel, which has several functions also in cutaneous biology,²⁹⁻³² reduces apoptotic features in an experimental skin model. This behavior is in accordance with previous studies, which had shown the down-regula-

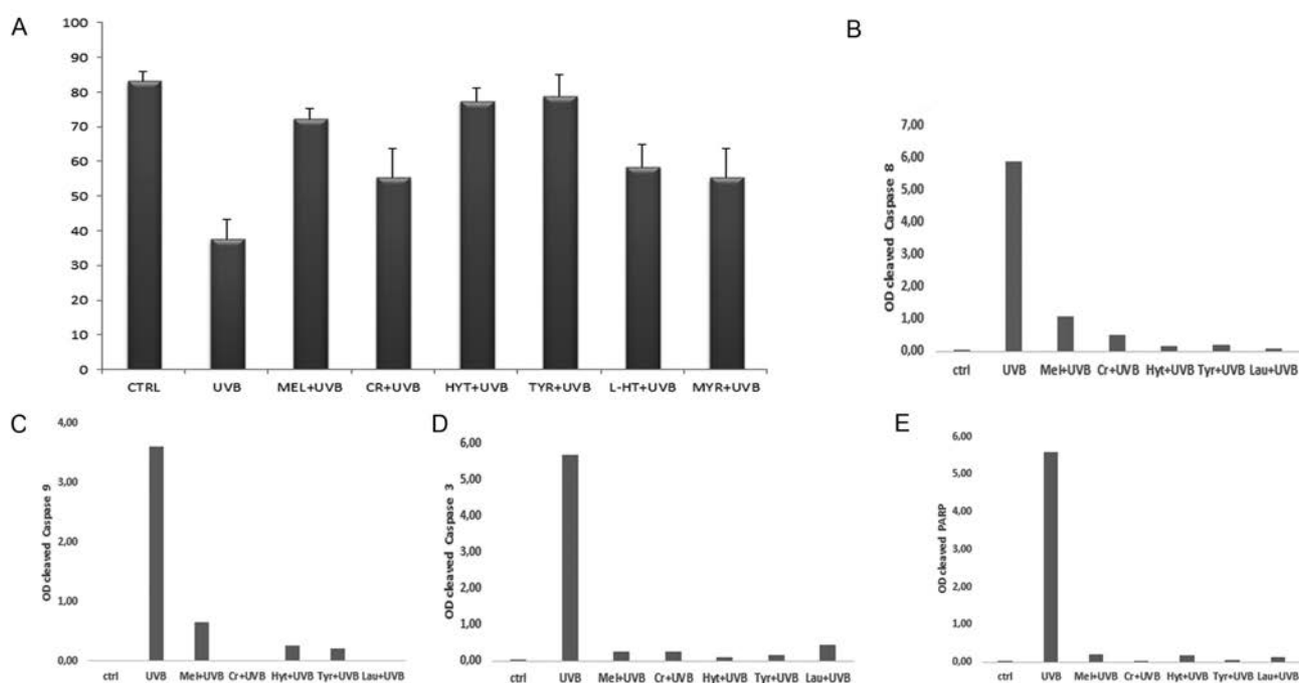


Figure 6. TB assay (A): percentage analysis of viable cells for each treatment. All pre-treatments reduced dead cell number in a significant way. Data are from three separate experiments and are presented as mean \pm standard deviation. Significance has been calculated through ANOVA test ($F=24.64$ with $P < 0.01$) and Tukey HSD; $**P < 0.01$ UVB *vs* CTRL, MEL, MEL, HYT, L-HT, TYR, MYR; $*P < 0.05$ UVB *vs* CR. Densitometric analysis of cleaved caspases-8, -9, -3 (B, C, D) and PARP(E) western blotting bands. Histograms evidenced no activation in untreated cells (ctrl). Caspases and PARP activation could be observed after UVB exposure while it was reduced by the different pre-treatments. Data were representative of three independent experiments performed in triplicate and SD was not shown because less than 10%. Data were expressed as ratio between mean optical density (OD) of cleaved caspase band and mean OD of actin band (protein control), in arbitrary units.

tion of mitochondrial cytochrome c release and the inhibition of caspase-9 and -3 activation in Mel treated samples.^{33,34} In addition, interestingly, human skin is able to produce Mel and its metabolites, with strong antioxidant activity^{35,36} exerting intriguingly inhibitory effects in melanoma cells³⁷ and suppressing activity of UVB-induced ROS formation.³⁸

On the other hand, Cr effect vs UVB radiation has been also demonstrated in keratinocytes *in vitro*. In particular, apoptotic signs, DNA fragmentation and caspase activation decrease, even if with a lesser extent, in Cr-treated samples before UVB exposure. Despite the important function of Cr as an essential energy precursor, its endogenous synthesis in humans, sometimes, is not sufficient and it has to be supplemented by daily food intake. Moreover, cutaneous cells may show a Cr level decrease, probably caused by stress or age-related dermal vascularization decline.³⁹ Thus, these findings suggest that Cr supplementation could result in a lower UVB-induced DNA damage in skin cells.

Finally, HyT and TyR, natural dietary polyphenols, well absorbed in the gastrointestinal tract, show anti-apoptotic effects against UVB radiation in HaCaT cells. HyT and TyR can be considered potential candidates to modify cellular response to UVB radiation. Moreover, these findings demonstrate that synthetic lipophilic hydroxytyrosyl esters can be considered anti apoptotic molecules able to significantly counteract UVB-induced damage, but with a lesser extent respect to the HyT. UVB radiation causes serious injury to skin cells and tissues (and especially to the epidermis) by inducing the production of inflammatory mediators and DNA lesions and, ultimately, by provoking cellular apoptosis.

In this work, the effect of different antioxidant compounds, chosen for their known antioxidant powers in different cell models, has been analyzed in human keratinocytes *in vitro*. These findings demonstrate that all compounds can be considered as anti-apoptotic agents against UVB-induced apoptosis. Antioxidants appear able to counteract the appearance of apoptotic ultrastructural features, such as chromatin condensation, cytoplasmic vacuolization and *in situ* DNA fragmentation, which clearly appear in cells exposed to UVB alone. Moreover, molecular analyses demonstrated the capacity of these compounds to prevent, at least in part, caspase activation, which were upregulated after UVB treatment alone.

This work also enhanced the role of oil polyphenol in preventing UVB apoptotic

cell death, evidencing the potential anti-apoptotic effect exerted by HyT or TyR, the main components of the phenolic fraction of virgin olive oil. Moreover, the effect of HyT esters of C12 or C14 fatty acids, with increased lipophilicity, has been evaluated to understand if the antiapoptotic effect also occurs in the presence of different number and position of the phenolic hydroxyl groups. Hyt derivatives with medium acyl side, such as lauryl or myristyl, are less efficient to protect keratinocytes against UVB than HYT. However, they are able to significantly counteract cell death and thank to their chemical structure, they penetrate, better than HyT, through the human corneum stratum and viable epidermis membranes as described by Procopio *et al.*⁴⁰

In conclusion, considered the general need for safe and effective skin protection against UVB-induced redox alterations, these compounds, able to reduce *in vitro* UVB-induced keratinocyte apoptosis, could represent important sun damage-preventing molecules, potentially useful *in vivo*, too.

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