Ultrasonographic assessment of the cutaneous changes induced by topical flavonoid therapy

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Abstract: Ultrasonography allows the quantification of dermal density and echogenicity changes during the physiological senescence process. Some active ingredients are able to slow down the tissue degeneration and disorganization process. The purpose of this study was to assess the cutaneous changes induced by the topical use of products containing Viniferol® as active ingredient, using high-frequency ultrasound. The study was performed over 12 weeks and included 80 healthy Caucasian female subjects, aged 22–75 years, divided into two groups: the study group and the control group. The product was applied according to a predetermined protocol. The measurements performed for each subject were: the thickness of the epidermis and dermis (mm), the number of low, medium, and high echogenic pixels, and the number of low echogenic pixels in the upper dermis/number of low echogenic pixels in the lower dermis. All the parameters showed a significant improvement. Ultrasound measurements showed an increase of the mean thickness of the epidermis \( P < 0.0001 \) and dermis \( P < 0.0001 \) following the application of the Viniferol product as compared to the control group. The changes in the dermal echogenicity confirm the efficacy and direct action of Viniferol upon the cutaneous fibroblasts. No side effects related to the treatment were recorded. The study proves the efficacy of this active ingredient in the cutaneous senescence process as well, as the fact that anti-aging prophylaxis should be initiated in the 20–40 year critical age group. This interval involves specific changes in dermal echogenicity that quantify intense molecular, biochemical and structural changes, being thus mostly and highly responsive to the anti-aging therapy.

Keywords: ultrasonography, flavonoid therapy, cutaneous changes, anti-aging therapy

Introduction

High-frequency ultrasound is a noninvasive diagnostic method with several applications in the field of internal medicine and, recently, clinical dermatology as well as dermatology and dermatocosmetics.1,2 It can represent a “noninvasive histological tool” for the quantification of the cutaneous structure or of the cutaneous pathology by monitoring the evolution of cutaneous lesions and the assessment of topical therapies’ efficacy in chronic inflammatory disorders (morphea, scleroderma, psoriasis), for the quantification of the intrinsic and extrinsic senescence process, as well as the efficacy of topical or generally applied therapies.3 It can also be used as an important research tool of the cutaneous senescence process and the efficacy of various anti-aging therapies.4,5

The use of high-frequency ultrasound in the assessment of the regenerative effects of various anti-aging therapies is an exciting and important area for research. From all the available imaging methods, high-frequency ultrasound (Dermascan 20 MHz Cortex Technology) is a new study tool focused on the assessment of the cutaneous
physiological and pathological features, having a penetration depth of up to 2.5 cm.

Collagen is an echogenic marker protein, synthesized by dermal fibroblasts, which can be identified by high-frequency ultrasound. The changes of the extracellular matrix, consisting in variations of the dermal density and echogenicity throughout the physiological senescence process, or during several anti-aging therapies quantify subtle reactions at molecular, cellular, and tissular level that influence the local homeostasis.8 High-frequency ultrasound allows an objective assessment of the cutaneous structure: the thickness of the epidermis and dermis as well as the dermal density, which are useful parameters for the quantification of the cutaneous regeneration process.7,8 Several changes of the cutaneous structure after topical prednisone therapy in children, or anti-aging natural supplement therapies in adults were ultrasonographically successfully assessed.9

Previous studies have shown that the thickness of the epidermis and dermis, as well as the dermal density are important parameters that assess the cutaneous regeneration process. The neosynthesis of the protein structures induces an increase of the dermal echogenicity and density, local cell architecture changes, and an increase of the dermis and epidermis thickness. The epidermis is the morphological expression of the changes in the subjacent dermis.7

It has been proved that certain ultrasonographic markers, such as SLEB (subepidermal low echogenicity band) or the LEPs/LEPi ratio (number of low echogenic pixels in the upper dermis/number of low echogenic pixels in the lower dermis) can quantify the cutaneous senescence process, as well as the efficacy of various anti-aging therapies.2,10,11

As far as the anti-aging therapy is concerned, Viniferol, a standardized extract from Bordeaux vine stalks, is one of the newest and very efficient anti-wrinkle and anti-aging agents. The active ingredient of Viniferol is ε-viniferine, a generic name for a new class of phytoalexins characterized as resveratrol dimers. Besides the well-known antioxidant effect of this flavonoid-containing extract, it has a direct action on the protein expression genes involved in the proliferation and differentiation of integumentary cells. These are retinoid-like effects which decrease cellular differentiation and increase cellular growth. Due to this action, Viniferol profoundly restructures and regenerates the skin and due to its antioxidant properties, it reestablishes the metabolic balance of the cutaneous cells, slowing down the tissular degeneration and disorganization process.12

Even though the general anti-aging effects of the flavonoids are well known, until now no scientific studies investigated the action of Viniferol at the cutaneous level by using high-frequency ultrasound.

The purpose of the study is the assessment, with the help of high-frequency ultrasound, of the cutaneous changes induced by topical use of products containing Viniferol.

**Materials and methods**

**Patients**

Eighty female subjects, aged 22–75 years, who requested prophylaxis and anti-aging therapy with flavonoids, were prospectively included in the study. Half (50%) of the subjects belonged to Fitzpatrick phototype class II and 50% to phototype class III.13 The study excluded patients with known allergies to topical flavonoids, cutaneous facial lesions, resurfacing or other anti-aging therapies in the last 2 months, or those who used phototherapy or oral contraception.

The subjects taken into the study were divided into two categories: a study group and a control group. The study group followed the proposed anti-aging therapy for 12 weeks, according to a standard protocol. In the morning, a hydrating emollient cream, based on occlusive hydrating agents was applied at the facial level (including the zygomatic area) by lightly massaging the area for 2 minutes.14 In the evening, an anti-aging product containing Viniferol, was applied in the same manner. No other cosmetic products were used by the subjects during the 12 weeks of study. The control group followed a placebo therapy for 12 weeks, using only moisturizing cream in the morning and evening, applied at the facial level, including the zygomatic area.

For every subject, ultrasonographic images were taken from the zygomatic level at the beginning and 12 weeks after the local application of the emollient, hydrating product, and the anti-aging, Viniferol-based cream.

The study was approved by the Ethical Committee of University of Medicine and Pharmacy Iuliu Hatieganu, Cluj-Napoca, Romania. Every subject was informed about the nature and purpose of the study, and signed a written consent before enrolling into the study.

**Ultrasonographic evaluation**

The ultrasonographic evaluation was performed with the Dermascan equipment (Cortex Technologies, Hadsund, Denmark), containing a 20 MHz transducer, that allows the “in vivo” acquirement of sectional cutaneous images up to a depth of 2.5 cm.

Dermascan consists of three major components: a transducer, an elaboration system, and a database (Figure 1).
The ultrasonic wave is partially reflected at the boundary between adjacent structures and generates echoes of different amplitudes. The intensity of the reflected echoes is evaluated by a microprocessor and visualized as a colored two-dimensional image.1 The color scale of echogenicity is: white – yellow – red – green – blue – black. Normally, the epidermal echogenicity appears as a white band, the dermis is expressed as a two-color composition: yellow and/or red, and the subcutaneous layer appears either green or black (Figure 2).

Statistical analysis
Statistical analysis was performed with SPSS (v 15.0; SPSS Inc, Chicago, IL). The data we obtained were analyzed, calculating the mean and standard deviation for the quantitative variables of every group and the proportions for the qualitative variables. The difference of means before and after treatment was tested using a t-test for paired samples, and the relationship between different parameters was assessed through Spearman’s correlation coefficients. A P value <0.05 was considered significant.

Results and discussion
Forty female subjects with a mean age of 49.60 ± 14.19 years (range 22–75 years) were included in the study group. Twelve subjects (30%) were under the age of 40, 16 subjects...
(40%) between 40–60 years and twelve (30%) over the age of 60. The control group was composed of subjects aged 22–75 years as well, divided into the above mentioned age categories.

All subjects involved in the study tolerated the therapy, without evoking adverse effects (erythema, pruritus, ocular disturbance). Subjectively, post-flavonoid therapy, a significant hydration of the skin throughout the day and an increase of the cutaneous tonicity was noticed.

The thickness of the epidermis in the study group initially was 0.129 ± 0.237 mm, and the thickness of the dermis 1.434 ± 0.241 mm.

After therapy, an increase of the mean thickness of the epidermis (0.129 ± 0.237 mm vs 0.150 ± 0.323 mm, \( P < 0.0001 \)), and of the dermis (1.434 ± 0.241 mm vs 1.569 ± 0.219 mm, \( P < 0.0001 \)) was observed (Tables 1 and 2).

The thickness of the dermis increased mainly in the 40–60 years age group (1.413 ± 0.280 mm vs 1.569 ± 0.279 mm, \( P = 0.001 \)), and less, but still significantly in the <40 years group (1.416 ± 0.266 mm vs 1.585 ± 0.150 mm, \( P = 0.015 \)), while in those >60 years, the increase was not statistically significant (1.480 ± 0.157 mm vs 1.554 ± 0.204 mm, \( P = 0.097 \)).

At the same time, at the dermal level, the number of LEP decreased (15,153.53 ± 3589.86 vs 12,958.48 ± 3628.35, \( P < 0.0001 \)), but this aspect was only noticed in the lower dermis (6949.75 ± 1966.93 vs 6257.62 ± 2224.88, \( P = 0.016 \)), not in the upper dermis (7290.55 ± 1794.60 vs 6940.65 ± 2150.30, \( P = 0.168 \)).

Overall, the LEPs/LEPi ratio increased significantly after flavonoid therapy (1.092 ± 0.330 vs 1.259 ± 0.631, \( P = 0.011 \)).

We also noticed an increase of MEP (3359.72 ± 1457.36 vs 3983.47 ± 1401.24, \( P = 0.013 \)) and HEP (460.27 ± 323.93 vs 750.90 ± 493.82, \( P < 0.0001 \)) after therapy.

The general variation pattern of the quantifiable ultrasonographic parameters after flavonoid therapy is illustrated in Table 1 and Figure 3.

If we consider the variation of the ultrasonographic parameters after topical flavonoid therapy according to the phototype class of the subjects, it can be noticed that after therapy, there is a significant increase of the LEPs/LEPi ratio in the subjects belonging to phototype class II, not III (Table 2).

In the placebo group, we noticed no significant thickening of the epidermis and a slight increase of the dermis after therapy (1.433 ± 0.34 mm vs 1.486 ± 0.14 mm). The number of LEP at dermal level also show a slight increase (13,213 ± 1284 vs 15,374 ± 2318, \( P = 0.1 \)) due to an optimal hydration of the skin and a discrete decrease of high echogenicity pixels (421.8 ± 121.18 vs 368.3 ± 104.03, \( P = 0.07 \)). The LEPs/LEPi ratio showed no particular display according to the age or phototype of the subjects.

The obtained results are in accordance with the data published in literature. Thus, locally applied flavonoids induce the neosynthesis of the fibrillary structures, but also of glycosaminoglycans, which display intense hydrophilic properties, favoring the cutaneous hydration.

It is well known that flavonoids have important anti-aging properties not only at a cutaneous level, but at the level of the entire organism. Viniferol, a molecule with proven anti-aging and antioxidant properties, exhibits a complex action at the cutaneous level: it interacts with fibroblastic receptors, amplifies the interrelation fibrocyte-extracellular matrix, modulates the adhesiveness molecules and interferes with the oxidative stress process and nonenzymatic glycation, with a regenerative effect at cutaneous level.

After therapy, a significant increase of the mean thickness of the epidermis and dermis was noticed, which once again confirms the presence of a complex, regenerative dermal process, induced by flavonoids.

The dermal thickness increased the most in the 40–60 years age group, to a lesser extent, but still significantly under the age of 40 years, and insignificantly over 60 years. We can affirm that topical flavonoid products have the best efficacy on mature integument, with specific structural and hormonal characteristics. In young subjects (<40 years) the thickness of the dermis increases discretely as the dermis is a young connective tissue, rich in glycosaminoglycans and thus, properly hydrated.

After the age of 60, the extracellular matrix (fibrocytes, fibers, glycosaminoglycans) of the skin undergoes characteristic degenerative changes. Therefore, the flavonoid-based anti-

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**Table 1** Cutaneous parameters quantified by high-frequency ultrasound before and after treatment

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Before treatment</th>
<th>After treatment</th>
<th>( P ) value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Thickness of epidermis (mm)</td>
<td>0.13 ± 0.24</td>
<td>0.15 ± 0.32</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>Thickness of dermis (mm)</td>
<td>1.43 ± 0.24</td>
<td>1.57 ± 0.22</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>LEP</td>
<td>15,153.53 ± 3589.86</td>
<td>12,958.48 ± 3628.35</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>MEP</td>
<td>3359.72 ± 1457.36</td>
<td>3983.47 ± 1401.24</td>
<td>0.013</td>
</tr>
<tr>
<td>HEP</td>
<td>460.27 ± 323.93</td>
<td>750.90 ± 493.82</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>LEPs</td>
<td>7290.55 ± 1794.60</td>
<td>6940.65 ± 2150.30</td>
<td>0.168</td>
</tr>
<tr>
<td>LEPI</td>
<td>6949.75 ± 1966.93</td>
<td>6257.62 ± 2224.88</td>
<td>0.016</td>
</tr>
<tr>
<td>LEPs/LEPi</td>
<td>1.09 ± 0.33</td>
<td>1.26 ± 0.63</td>
<td>0.011</td>
</tr>
</tbody>
</table>
Table 2 Variation of the cutaneous parameters quantified by high-frequency ultrasound before and after treatment, according to phototype

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Prototype III Before treatment</th>
<th>Prototype III After treatment</th>
<th>P value</th>
<th>Prototype II Before treatment</th>
<th>Prototype II After treatment</th>
<th>P value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Epidermis (mm)</td>
<td>0.1288 ± 0.026</td>
<td>0.151 ± 0.027</td>
<td>&lt;0.0001</td>
<td>0.129 ± 0.021</td>
<td>0.148 ± 0.037</td>
<td>0.020</td>
</tr>
<tr>
<td>Dermis (mm)</td>
<td>1.441 ± 0.270</td>
<td>1.570 ± 0.263</td>
<td>0.001</td>
<td>1.427 ± 0.214</td>
<td>1.568 ± 0.172</td>
<td>0.003</td>
</tr>
<tr>
<td>LEP</td>
<td>15.059 ± 4063.97</td>
<td>13.864 ± 3824.33</td>
<td>0.015</td>
<td>15.247 ± 3162.14</td>
<td>12.052 ± 730.34</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>MEP</td>
<td>3120.10 ± 1725.95</td>
<td>3850.95 ± 1487.32</td>
<td>0.046</td>
<td>3599.35 ± 1122.42</td>
<td>4116.0 ± 1334.62</td>
<td>0.151</td>
</tr>
<tr>
<td>HEP</td>
<td>379.35 ± 280.94</td>
<td>645.50 ± 373.59</td>
<td>&lt;0.0001</td>
<td>541.20 ± 350.25</td>
<td>856.3 ± 581.03</td>
<td>0.004</td>
</tr>
<tr>
<td>LEPs</td>
<td>7071.65 ± 1754.22</td>
<td>6961.1 ± 2236.66</td>
<td>0.725</td>
<td>7509.45 ± 1852.70</td>
<td>6920.20 ± 2118.53</td>
<td>0.148</td>
</tr>
<tr>
<td>LEPI</td>
<td>7007.65 ± 2150.80</td>
<td>6737.15 ± 2205.91</td>
<td>0.480</td>
<td>6891.85 ± 1818.86</td>
<td>5778.10 ± 2193.29</td>
<td>0.010</td>
</tr>
<tr>
<td>LEPs_LEPi</td>
<td>1.035 ± 0.262</td>
<td>1.096 ± 0.300</td>
<td>0.200</td>
<td>1.1499 ± 0.384</td>
<td>1.4227 ± 0.820</td>
<td>0.026</td>
</tr>
</tbody>
</table>

Aging therapy has less effective regenerative changes in this age interval. These regenerative changes could be amplified through the use of products able to interfere with the characteristic aging mechanisms for this specific age group.

At the same time, concomitantly with the change in dermal thickness, the number of LEP decreased in the lower dermis, but not in the upper dermis. The LEPs/LEPi ratio also increased significantly after therapy. The decrease in the number of LEP in the lower dermis is proportional to the significant increase of MEP and HEP that quantify protein neosynthesis, as well as cytoarchitectural reorganizations of the extracellular matrix.

According to the literature, the cutaneous imaging parameters that can be considered as objective in the evaluation of the cutaneous aging/regeneration process are the following:

- **LEP** quantifies the degree of cutaneous hydration, inflammatory processes, solar elastosis, collagen degeneration.
- **MEP** and **HEP** quantify the structures of collagen and elastin fibers and microfibrils.
- The LEPs/LEPi ratio allows an appreciation of the density and integrity of the extracellular matrix, both from the upper and lower dermis, which vary according to age, UV-ray exposure, and therapy.16

The data obtained from the study shows important ultrasonographic changes at the cutaneous level after anti-aging therapy. Flavonoids have a complex action at the dermal level, interfering with several mechanisms involved in the senescence process. They act at the level of fibrocytes, on specific receptors, turning inactive mature cells into young, metabolically active ones.19

Also, studies performed in order to assess the phyto-oestrogen activity of Viniferol (evaluation of gene expression, coding for typical differentiation proteins) have shown that it influences gene expression and cell proliferation both at the epidermal and dermal levels. Therefore Viniferol decreases gene expression in *Loricrine, Fillagrin, and Keratin 11*, and increases gene expression in *Calgranulin B*.20

The synthesis of the protein structures is initiated at the intracellular level. The trophocollagen molecules, elastin, and the glycosaminoglycans are extracellularly assembled into microfibrils, fibers, or proteoglycans. Depending on the biochemical structure, the level of organization, architectural orientation, and quantity, the proteins show a certain cutaneous echogenity degree. The low echogenity pixels that quantify the hydration degree of the extracellular matrix especially in the lower dermis, are replaced by medium and high echogenity pixels, quantifying protein synthesis. The MEP codify the presence of elastin and collagen microfibrils. Once these are assembled into mature fibers, the local echogenicity increases, and are expressed as hyperechogenic areas (HEP). The increase of the LEPs/LEPi ratio quantifies the replacement of the hypoechogenic pixels from the lower dermis with medium and high echogenic pixels as a result of protein neosynthesis. Type I collagen, that is predominant at the dermal level (punctiform hyperechogenic pixels) is organized in fibers, visible as hyperechogenic bands, having a parallel
display in the lower dermis. These hyperechogenic bands, visible especially on photoprotected sites, represent an ultrasonographic marker of the intrinsic aging process.\textsuperscript{21}

The collagen–elastin protein structure, in different organization stages, is quantified by pixels of different amplitude. The post-therapeutic changes, visible on the sonograms as an increase of medium and high echogenic pixels, are suggestive of increased protein synthesis, the assembly of molecules to filaments, fibrils, and eventually, collagen fibers. According to our observations, there is a direct relationship between the genetically determined cutaneous phototype, the structure and repartition of echogenic proteins (collagen, elastin) and the different response to anti-aging therapies.\textsuperscript{22}

If we consider the significant changes of the ultrasonographic parameters after anti-aging therapy depending on the phototype of the subject, a significant increase of the LEPs/LEPi ratio is present in the subjects in phototype II class, but not class III. This observation would justify the correlation of the anti-aging therapy with the cutaneous phototype. Further studies are necessary to confirm the different reactivity of the phototype classes to local therapies.

The synthesis process of fibrillary proteins depends on several factors, involving complex, targeted, and personalized therapies. According to the “wear and tear” theory, throughout life, intrinsic and extrinsic mechanisms associated to the senescence process reduce the regenerative capacity of cells, leading to post-translational changes at the level of intracellular and extracellular proteins, such as oxidation and nonenzymatic glycation.

Flavonoids, through complex mechanisms, interfere with the reactions involved in the senescence process, and induce the synthesis of the extracellular matrix. According to our data, Viniferol-based products are more efficient in the 40–60 years age group, characterized by complex biological changes at the cutaneous level. Viniferol shows real and important anti-aging properties, since it interferes concomitantly with the genetic, oxidative, immunologic, and metabolic mechanisms that are involved in the cutaneous aging process.\textsuperscript{23}

Prophylaxis of the aging process should start before the age of 40 years, preferably in the critical age group (20–40 years), which is characterized by important changes at the tissue, cellular, and molecular levels.\textsuperscript{24} The optimization of anti-aging therapy, according to special studies, requires targeted, personalized therapies, adapted to the hormonal, genetic, oxidative, immunologic, and metabolic status of the subject, and is capable of interfering with deficient mechanisms in certain age groups.

High-frequency ultrasound can be used as a modern, noninvasive, histological tool for the assessment of various anti-aging therapies, with applications in the fields of pharmacology and dermatocosmetology.

Conclusions

High-frequency ultrasound is a modern, noninvasive method that allows the assessment of the efficacy of anti-aging therapies. After topical flavonoid-based therapy, a significant increase of the epidermis and dermis was noticed, especially in the 40–60 years age group. Concomitantly, other cutaneous changes take place: decrease in the number of LEP and increase in the MEP and HEP. The cutaneous changes are more marked in subjects belonging to phototype class II. Anti-aging prophylaxis should be initiated in the 20–40 years critical age group, involving personalized therapies also adapted to the phototype.

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Disclosure

The authors report no conflicts of interest in this work.

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