**In vitro** induction of melanin synthesis and extrusion by tamoxifen

T. Matama*, R. Araújo*, A. Preto*, A. Cavaco-Paulo† and A. C. Gomes*

*CBMA (Centre of Molecular and Environmental Biology), Department of Biology, University of Minho, Campus of Gualtar, 4710-057, Braga, Portugal and †IBB – Institute for Biotechnology and Bioengineering, Centre of Biological Engineering, University of Minho, Campus de Gualtar, 4710-057, Braga, Portugal

Received 01 February 2013, Accepted 06 April 2013

**Keywords:** catalase, cell culture, colour cosmetics, genetic analysis, SERM, skin pigmentation

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**Synopsis.**

**OBJECTIVES:** Physical appearance has significant importance psychologically and socially, with skin and hair being of prime relevance. Effective ingredients that modulate melanin synthesis are of growing interest. Tamoxifen, a widely used selective oestrogen receptor modulator, SERM, was described occasionally in medical case reports as causing grey hair repigmentation. This work aimed to study, *in vitro*, the effect of tamoxifen and 4-hydroxy-tamoxifen, one of its most bioactive derivatives, on melanin production in human melanocytes.

**METHODS:** Adult normal human epidermal melanocytes (NHEM) were treated with physiological concentrations of tamoxifen and 4-hydroxy-tamoxifen during 72 hours. Cytotoxicity was evaluated by lactate dehydrogenase (LDH) leakage. Total melanin was quantified by spectrophotometry, and cyclic adenosine monophosphate (cAMP) was determined by competitive ELISA. The relative mRNA levels of several genes involved in melanogenesis were investigated by real-time PCR.

**RESULTS:** Under the conditions used, the results showed that tamoxifen and 4-hydroxy-tamoxifen treatments, none of them toxic to NHEM, induced a time-dependent increase in the amount of melanin released to the culture medium. cAMP, one of the major second messenger in signalling pathways important to melanogenesis, was decreased after treatment. The transcript levels of genes coding for catalase, premelanosome protein and melan-A, directly related to skin and hair pigmentation, showed an increased tendency upon tamoxifen and 4-hydroxy-tamoxifen treatment. Induction of catalase gene expression in NHEM points towards a promelanogenic effect mediated by ROS.

**CONCLUSION:** According to the results, even in such a short treatment period, tamoxifen and 4-hydroxy-tamoxifen promoted melanin extrusion and they seem to act as melanogenesis stimulators at the molecular level. Our data suggest that SERMs might be a new tool for increasing melanogenesis and might be of great interest for topical formulations in cosmetic industry.

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**Résultat.**

**OBJECTIF:** L’apparence physique a une grande importance psychologique et sociale, la peau et les cheveux étant d’un intérêt primordial. Les ingrédients efficaces qui modulent la synthèse de mélanine sont d’un intérêt croissant. Le tamoxifène, un modulateur sélectif des récepteurs d’oestrogène largement utilisé (SERM), a été décrit dans les rapports de cas médicaux comme causant une ré-pigmentation des cheveux gris. Ce travail visait à étudier, *in vitro*, l’effet du tamoxifène et du 4-hydroxy-tamoxifène, un de ses dérivés les plus bioactifs, sur la production de mélanine dans les mélanocytes de l’homme.

**MÉTHODES:** Des mélanocytes épidermiques adultes humains normaux (NHEM) ont été traités avec des concentrations physiologiques de tamoxifène et de 4-hydroxy-tamoxifène pendant 72 heures. La cytotoxicité a été évaluée par la sécrétion de la lactate-déshydrogénase (LDH). La mélanine totale a été quantifiée par spectrophotométrie et l’adenosine monophosphate cyclique (AMPc) a été déterminée par ELISA. Les niveaux relatifs d’ARNm de plusieurs gènes impliqués dans la mélanogénèse ont été étudiés par PCR en temps réel.

**RÉSULTATS:** Dans les conditions utilisées, les résultats ont montré que les traitements au tamoxifène et au 4-hydroxy-tamoxifène, aucun d’entre eux n’étant toxique pour les NHEM, induisent une augmentation en fonction du temps, de la quantité de mélanine libérée dans le milieu de culture. L’AMPc, un des messagers secondaires importants dans les voies de signalisation principales de la mélanogénèse, a diminué après le traitement. Les niveaux de transcription des gènes codant pour la catalase, le premelanosome et melan-A, directement liés à la pigmentation de la peau et des cheveux, ont montré une tendance à l’augmentation par le tamoxifène et le 4-hydroxy-tamoxifène. L’induction de l’expression du gène de la catalase dans les NHEM pointe vers un effet pro-mélanogénique médial par les ROS.

**CONCLUSION:** Selon les résultats, même dans une période de traitement de courte durée, le tamoxifène et le 4-hydroxy-tamoxifène promeuvent l’extrusion de la mélanine et ils semblent agir comme stimulateurs de la mélanogénèse au niveau moléculaire. Nos données suggèrent que les SERMs pourraient devenir un nouvel outil pour augmenter la mélanogénèse et peut-être avoir un grand intérêt pour les formulations topiques dans l’industrie cosmétique.

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**Introduction**

In today’s society, the concern with preserving a good physical appearance feeds a huge industry of cosmetic products, with skin and its appendages (hair in particular) being the primary targets.
Oestrogens play important protective roles in skin physiology and ageing [1]. They accelerate wound healing, improve inflammatory skin conditions and provide protection to skin photoaging [2]. The beneficial effects are confirmed by the changes seen in postmenopausal women when a rapid skin age-related deterioration is observed – thinner and drier skin, more and deeper wrinkles, loss of skin firmness and elasticity [2]. Topical application of oestrogens and hormonal replacement therapies can reverse at some extent these skin age-related events. The effect of oestrogens in human hair cycle is not so obvious; they appear to stimulate hair growth by decreasing the resting phase and by prolonging the anagen phase, especially in women; it was demonstrated that oestrogen has the ability to modify androgen metabolism in dermal papillae of hair follicles [3]. Oestrogens also affect skin pigmentation. High oestrogen levels are associated with an increase in skin pigmentation particularly evident during pregnancy and during the use of oestrogen-containing oral contraceptives. Topical application of oestrogens was described as having no significant effect on pigmentary changes in aged skin [4]. In *in vitro* studies, some apparent conflicting reports have been published; McLeod et al. [5] reported that melanocytes respond to oestrogens, in particular to 17β-oestradiol, with increased melanin production and extrusion, whereas Jee et al. [6] reported an increase in melanocyte proliferation with a concomitant decrease in melanin levels and tyrosinase activity. More, recently, it was demonstrated in *in vitro* that the oestrogen receptor β is an activator of dopachrome tautomerase gene (DCT) promoter activity in cooperation with additional transcription factors and microphthalmia-associated transcription factor (MITF) in normal human melanocytes [7].

Since the discovery in the 1990s that skin expresses oestrogen receptors that decline in number from the onset of menopause, the search for sale and effective alternatives with more focused effects on the skin, topical oestrogens, phyto-oestrogens and tissue-specific drugs called selective oestrogen receptor modulators (SERMs) has been explored [8, 9]. SERMs bind to oestrogen receptors and can act as either oestrogen agonists or antagonists depending on the target tissue. Tamoxifen, a non-steroidal triphenylethylene, is a SERM widely used, since 1971. In the management of women with all stages hormone-responsive breast cancer [10]. Although the effects of oestrogens on skin are extensively documented, there are very limited data on the effect of SERMs on this organ. Topical tamoxifen improves the appearance of keloid scars in acute burns patients, fading fibroblast proliferation and collagen synthesis [1]. SERMs tamoxifen, raloxifene and genistein all substantially benefit skin healing in the oestrogen-deprived ovarioctomized mouse, most likely mediated via oestrogen receptor β [1]. In contrast, a subcutaneous administration of tamoxifen in rat resulted in the appearance of abnormal hair follicles, epidermal atrophy and increased dermal fibrosis, particularly around the hair follicles; in humans, there have been reports of tamoxifen treatment causing diffuse thinning of the hair with moderate receding of the frontal hair line and the development of alopecia on the crown; in organ culture, tamoxifen alone has no effect on human hair shaft elongation, whereas in excess tamoxifen in combination with 17β-oestradiol eliminated the effect of 17β-oestradiol, suggesting that tamoxifen can function as an antagonist of oestrogen in the scalp hair follicle [2]. Tamoxifen has also been described as causing repigmentation of a 68-year-old breast cancer patient’s grey hair [11]. So far, the available literature points to the ability of SERMs acting as oestrogen agonists in skin, being able to control skin ageing, wound healing and scarring and to their potential use in topical formulations that would avoid the harmful effects associated with systemic hormone replacement therapies [8].

Along with the beneficial effects of SERMs already described on skin healing and ageing, it would be of great interest to ascertain their effects on skin pigmentation. Melanogenesis is a defensive mechanism against excessive exposure of human skin to sunlight, an environmental risk factor for melanoma and skin ageing. This work aims to study, *in vitro*, the effect on melanin production of tamoxifen and 4-hydroxy-tamoxifen, one of its most bioactive derivatives and to contribute to a more complete understanding of the potential cosmetic use of this SERM.

**Materials and methods**

**Reagents**

Tamoxifen (Tam) and 4-hydroxy-tamoxifen (4HO-Tam), reduced disodium salt hydrate β-nicotinamide adenine dinucleotide (NADH), sodium pyruvate, melanin from *Sepia officinalis* and Trypsin–EDTA solution were purchased from Sigma-Aldrich Chemie GmbH (Buchs, Switzerland). Soluene-350 was obtained from PerkinElmer (Waltham, MA, U.S.A.). Absolute ethanol (BIOH) used as solvent for Tam and 4HO-Tam, as well as other common chemical reagents used to prepare the buffers, was purchased from Panreac (Barcelona, Spain).

**Cells and culture conditions**

Adult normal human epidermal melanocytes (NHEM) and the optimized media for their growth were supplied by Lonza (Lonza Walkersville, Inc, Walkersville, MD, U.S.A.). Cells were maintained in Melanocyte Cell Basal Medium-4 supplemented with calcium chloride, bovine pituitary extract, recombinant human fibroblast growth factor-B, recombinant human insulin, hydrocortisone, phorbol 12-myristate 13-acetate, gentamicin sulphate and amphotericin-B, foetal bovine serum (melanocyte growth medium 4, MCM4) and endothelin-3 according to the supplier’s instructions. Cell culture was established and maintained in 25 cm² flasks at 37°C and 5% CO₂ in a humidified incubator with the recommended starting cell density of 10 000 cells cm⁻².

**NHEM treatment with tamoxifen and 4-hydroxy-tamoxifen**

Normal human epidermal melanocytes were treated with physiological concentrations of the drugs. Two days after seeding passage 3, melanocytes were incubated with 1 μM Tam and 0.5 μM 4HO-Tam for 72 h. As controls, melanocytes without any treatment and melanocytes incubated with 0.1% BIOC were plated in parallel. The media was refreshed every 24 h and collected for further analysis.

**Total melanin spectrophotometric evaluation**

The melanin content was quantified with either 300 000 cells, plated on Petri dishes (2560 mm), or from the culture media collected at 24 h, 48 h and 72 h. For measuring intracellular melanin, cells were detached and disrupted in one millilitre of 15 mM Tris-HCl lysis solution, pH 7.5. A 150 μl of either cell lysate or cell culture medium was solubilized in 1350 μl of Soluene-350 at 80°C for 1 h. The calibration curves were obtained by solubilizing 5, 10, 20, 30, 40, 60, 80, 100, 300 and 500 mg of *Sepia* melanin in Soluene-350 with 10% Tris-HCl or culture medium, depending on sample type, at 80°C for 1 h [12, 13]. Before measuring the absor-
bance at 500 nm, samples and standards were centrifuged for 10 min at 8600 g.

**Cytotoxicity evaluation**

The cytotoxicity induced by Tam and 4HO-Tam at the tested concentrations was evaluated by lactate dehydrogenase (LDH) release to the culture medium simultaneously to extruded melanin monitoring. The LDH activity was determined by the decrease in NADH absorbance at 340 nm during 3 min at 30°C, using a microplate reader. The final reaction volume was 300 μL consisting in 250 μM NADH and 280 μM pyruvate in 50 mM phosphate buffer, pH 7.4. To measure the intra- and extracellular LDH activity, 45 μL of NHEM lysate and 90 μL of culture medium were used, respectively. The percentage of LDH release was expressed as the relation between extracellular activity and total LDH activity.

**Determination of cyclic adenosine monophosphate levels**

To assess how NHEM treatment with Tam and 4HO-Tam affects the levels of cytoplasmic cyclic adenosine monophosphate (cAMP), 145 000 cells were seeded on 6-well plates and treated as described. After 72 h, cells were collected in 300 μL of 100 mM HCl solution. Cyclic AMP was quantified using a commercial competitive enzyme-linked immunosorbent assay (Cayman Chemical, Ann Arbor, MI, U.S.A.).

**Relative gene expression levels determined by real-time PCR**

Total RNA was isolated from 55 500 melanocytes, cultured in 12-well plates and treated with Tam and 4HO-Tam as described before, using the SV Total RNA Isolation System (Promega Corporation, Fitchburg WI, U.S.A.) according to the manufacturer’s instructions. Reverse transcription (RT) was performed with iScript cDNA Synthesis Kit (Bio-Rad Laboratories, Lda., Amadora, Portugal), according to the supplied protocol. The reaction was carried out at 25°C for 5 min, 42°C for 60 min and 85°C for 5 min. Real-time PCR was used to determine whether Tam/4HO-Tam were able to affect the expression of catalase gene and several genes overexpressed in human pigmented hair follicles when compared with grey hair follicles (unpublished results). TaqMan Universal Master Mix II and the pre-designed/pre-formulated primer and probe sets, TaqMan Gene Expression Assays (Table I), were purchased from Applied Biosystems (Foster City, CA, U.S.A.), and the ΔΔCt method was applied using as housekeeping gene, the mitochondrial ATPase 6 (MT-ATP6). The PCR amplification was performed according to the manufacturer’s instructions on an CFX96 Touch Real-Time PCR Detection System (Bio-Rad Laboratories, Lda.).

**Statistical analysis**

The data were expressed as the mean ± SEM of at least five replicates. A nonparametric analysis was performed: Kruskal–Wallis H-test or Friedman test for repeated measures along time. When P-values were equal or inferior to 0.05, the differences were considered significant. Whenever the Kruskal–Wallis H-test led to significant results, Mann–Whitney U-test was performed for sample contrasts between individual sample sets at 0.05 level of risk. All the statistical analysis was performed using Minitab 16 version 16.1.0 (Minitab Inc., State College, PA, U.S.A.).

**Results and discussion**

In the present work, we studied the effect of tamoxifen and 4-hydroxy-tamoxifen on melanin production. For this purpose, normal human epidermal melanocytes (NHEM) were incubated with 0.1% (v/v) ethanol solutions of tamoxifen (Tam) and 4-hydroxy-tamoxifen (4HO-Tam) at concentrations in physiological range as determined in patients’ serum under tamoxifen treatment [14]. The cells were exposed during three days with medium renewal every 24 h. Total melanin was quantified by spectrophotometry. After performing a calibration curve, the average intracellular melanin content of control NT after 72 h was determined as 540 ng μg⁻¹ of protein. The obtained results were expressed as a ratio of normalized melanin amount having the control NT as divisor (Fig. 1). A trend was observed in the amount of intracellular melanin; melanocytes exposed to Tam and 4HO-Tam produced significantly more melanin than the controls, although ethanol alone (Control

<table>
<thead>
<tr>
<th>TaqMan assay</th>
<th>Gene symbol</th>
<th>Protein name</th>
<th>Function (GeneCards)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hs00156308_m1</td>
<td>CAT</td>
<td>Catalase</td>
<td>Protection of cells from the toxic effects of hydrogen peroxide</td>
</tr>
<tr>
<td>Hs00167051_m1</td>
<td>TYRP1</td>
<td>Tyrosinase-related protein 1</td>
<td>Melanosomal enzyme that belongs to the tyrosinase family and plays an important role in the melanin biosynthetic pathway</td>
</tr>
<tr>
<td>Hs00173854_m1</td>
<td>SILV</td>
<td>Premelanosome protein</td>
<td>Melanocyte-specific type I transmembrane glycoprotein (PMEL17) involved in the maturation of melanosomes from stage I to II</td>
</tr>
<tr>
<td>Hs00194133_m1</td>
<td>MLANA</td>
<td>Melan-A (MART-1)</td>
<td>Vital role in the expression, stability, trafficking and processing of melanocyte protein PMEL17</td>
</tr>
<tr>
<td>Hs01385410_m1</td>
<td>SLC24A5</td>
<td>Solute carrier family 24, member 5</td>
<td>Cation exchanger involved in pigmentation, possibly by participating in ion transport in melanosomes</td>
</tr>
<tr>
<td>Hs01099965_m1</td>
<td>TYR</td>
<td>Tyrosinase</td>
<td>Copper-containing oxidase that functions in the formation of pigments such as melanins and other polyphenolic compounds</td>
</tr>
<tr>
<td>Hs00174029_m1</td>
<td>KIT</td>
<td>v-kit Hardy-Zuckerman 4 feline sarcoma viral oncogene homolog</td>
<td>Type 3 transmembrane receptor for MGF (mast cell growth factor, also known as stem cell factor)</td>
</tr>
<tr>
<td>Hs00955621_m1</td>
<td>PLXNC1</td>
<td>Plexin C1</td>
<td>Mitochondriaally encoded ATP synthase 6</td>
</tr>
<tr>
<td>Hs02596862_g1</td>
<td>MT-ATP6</td>
<td></td>
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</tbody>
</table>
EtOH) was able to induce melanogenesis, to a smaller extent, under this experimental conditions. In a genome-wide siRNA-based functional genomics study of pigmentation, unexpected genes were found to support pigment production through tyrosinase expression and stability [15]. Two of those genes encode two isoforms of aldehyde dehydrogenase, ALDH1A1 and ALDH9A1, which regulate ethanol detoxification. Non-toxic chemical inhibitors of the two ALDH enzymes inhibited pigmentation and tyrosinase protein accumulation in MNT-1 cells, and they impaired UV-induced tyrosinase expression when tested in primary melanocytes [15]. On the other hand, the amount of ethanol solvent present in cell culture medium could promote the activity of these detoxifying enzymes and promote melanogenesis in NHEM (Fig. 1).

Skin pigmentation is the result of two major events: the synthesis of melanin by melanocytes and the transfer of melanosomes to the surrounding keratinocytes [16]. Thus, for its biological relevance, melanin extrusion was also evaluated during the incubation of NHEM with Tam and 4HO-Tam (Fig. 2). The results obtained revealed a significant kinetic effect of Tam and mainly 4HO-Tam on melanin extrusion by NHEM. Note that 4HO-Tam accelerated melanin extrusion in the first 24 h, although these levels became similar in all conditions at 48 h and 72 h.

In normal human cultured melanocytes, physiological concentrations of oestrogens, especially 17β-oestradiol, were described as enhancers of melanin production (1.25-fold increase) and extrusion (3.4-fold increase after 16 h) [5]. The authors also tested the steroid anti-oestrogen ICI 164 384, and they verified that in melanocytes, it also increased tyrosinase activity, behaving as an oestrogen agonist. In the experiment performed, it becomes apparent that Tam and particularly 4HO-Tam behave as oestrogen agonists in NHEM by promoting both melanogenesis and melanin extrusion. The fact that the MGM-C226 medium was supplemented with endothelin-3 to enhance the proliferative capacity of melanocyte cultures through serial passages, according to manufacturer instructions, may have diluted the additive response of cells to Tam and 4HO-Tam because the basal levels of control were raised by this hormone (data not shown) [17]. On the other hand, the pro-melanogenesis effect of tamoxifen may be a long-term effect as reported for the repigmentation of a 68-year-old breast cancer patient’s grey hair after two years of tamoxifen treatment [11].

Cell viability was monitored by evaluating cell membrane permeability. The amount of lactate dehydrogenase (LDH) released into the medium was determined in terms of its enzymatic activity and related to the activity in the medium of non-treated (NT) NHEM, after the first 24 h of incubations (Fig. 3). In all the treatments, the LDH release was under 5% during the assay period. As expected, the amount of LDH leakage revealed that Tam and 4HO-Tam had no significant cytotoxic effect at the tested concentrations. Another control was performed bearing the same solvent concentration.
tration present in Tam and 4HO-Tam treatments; this control (Control EtOH) also did not affect the NHEM viability (Fig. 3).

Oestrogens regulate biological functions by two distinct mechanisms: a classical pathway that involves the binding to nuclear receptors (ligand-inducible transcription factors) and consequent regulation of the expression of target genes and a non-classical pathway that begins with the oestrogen binding to a membrane receptor followed by rapid intracellular signalling cascades that can eventually lead to gene transcription regulation. In normal human cutaneous melanocytes, the presence of classical oestrogen receptors has been demonstrated by binding studies and using immunocytochemistry and RT-PCR [6, 18]. Although SERMs may act through alternative mechanisms not involving the oestrogen receptor binding, to better elucidate their biological impact, the transcript level of genes of interest should be monitored [19, 20]. The expression of melanogenesis-related genes is crucial to the synthesis of pigments. Hence, in this work, the outcome after 72 h of Tam and 4HO-Tam treatment on transcription of melanogenesis-related genes in NHEM was evaluated by real-time PCR (Fig. 4). These genes were identified in a microarray study by us (unpublished) as being the most expressed in pigmented vs. grey hair follicles. In NHEM, after 72 h of treatment, 4HO-Tam in particular up-regulated melanogenesis-related genes transcription even after melanin extrusion peak at 24 h. Even so, the fold change in TYR mRNA induced by 4HO-Tam is in the range described by Kippenberger et al. [21], after treating normal human melanocytes with 20 μM oestradiol for 48 h. Considering the control 0.1% ethanol (control EtOH), the effect of 4HO-Tam and, to a lesser extent, the effect of Tam were more pronounced for SILV and MLANA genes. PMEL17, also known as gp100 or SILV, is a structural protein essential for the transition from stage I to stage II melanosomes. After its delivery to stage I melanosomes, PMEL17 is cleaved into several fragments, which form the fibrillar matrix of the organelle [22]. In pigmented cells, melamins are deposited on these protein fibres, resulting in a progressively pigmented internal matrix. MART-1, codified by MLANA gene, forms a complex with PMEL17 and affects its post-transcriptional production, stability, trafficking and processing [23]. 4HO-Tam and Tam affect in particular these two genes involved in the early maturation of melanosomes. Despite the absence of known diseases associated with these genes, it is possible that some forms of albinism may be linked to PMEL17 deficiency by analogy to the mouse silver mutant [24]. Eventually, these SERMs may be a good starting point for future investigation on therapeutic treatment in heterozygosis disorders involving PMEL17 or MART-1.

To further understand the possible mechanism of action of 4HO-Tam and Tam in NHEM, the levels of cAMP, a key messenger in melanogenesis, were determined (Fig. 5). The solvent and both Tam and 4HO-Tam significantly decreased the levels of intracellular cAMP (Fig. 5). Levels of intracellular cyclic adenosine monophosphate (cAMP) in NHEM after 72 h of incubation with tamoxifen (Tam) and 4-hydroxy-tamoxifen (4HO-Tam). The effect of the solvent alone (control EtOH) is also shown. Averages that do not share a letter are significantly different according to Kruskal–Wallis H-test (H = 22.48; DF = 3; P < 0.001) followed by Mann–Whitney U-test as post-hoc test.

Figure 4 Relative expression of melanogenesis-related genes normalized to transcript levels of housekeeping gene mitochondrial ATPase 6 in NHEM after 72 h of incubation with tamoxifen (Tam) and 4-hydroxy-tamoxifen (4HO-Tam), determined by real-time PCR using the ΔΔCt method. The effect of the solvent alone (control EtOH) is also shown. No significant differences were detected by the Kruskal–Wallis H-test.

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International Journal of Cosmetic Science, 1–7

Figure 6 Relative gene expression of catalase normalized to transcript levels of housekeeping gene mitochondrial ATPase 6 in NHEM after 72 h of incubation with tamoxifen (Tam) and 4-hydroxy-tamoxifen (4HO-Tam), determined by real-time PCR using the ΔΔCt method. The effect of the solvent alone (control EtOH) is also shown. Averages that do not share a letter are significantly different according to Kruskal–Wallis H-test (H = 6.51; DF = 2; P = 0.039) followed by Mann–Whitney U-test as post-hoc test.
lar cAMP. The results have shown that after 72 h, the levels of cAMP are negatively correlated with the levels of intracellular melanin, indicating that the chosen time window, although revealing significant changes in melanin content, did not support the notion that the mechanism of action involves elevation of cAMP levels. In melanocytes, the cAMP/CREB signalling pathway is regulated by melanocyte-stimulating hormone via the G-protein-coupled receptor MC1R, which in turn transcriptionally activates expression of MITF, the master transcriptional regulator of melanocyte development [25]. In vitro and in vivo, the up-regulation of cAMP levels in melanocytes leads to increased pigment production. The magnitude and duration of cAMP action are regulated by both production and hydrolysis. MITF directly regulates the transcription of the phosphodiesterase 4D3, creating negative homeostatic control of the cAMP pathway to counteract chronic stimulation of the cAMP pathway in melanocytes [26]. Fast response to oestrogens is attributed to membrane oestrogen receptors that trigger rapid increases in cAMP, calcium, extracellular signal-regulated kinase (ERK), signalling pathways all involved in melanogenesis regulation [27]. Tamoxifen and 4HO-Tam are able to bind to G-coupled membrane oestrogen receptor possessing also a non-genomic mechanism of action as oestrogen agonists [28–31].

Melanin synthesis generates hydrogen peroxide (H₂O₂) and other free radicals making oxidative stress a common theme in melanocyte dysfunction especially relevant for the hair follicle melanocyte population [32]. It was shown in vivo that human grey/white scalp hair shafts accumulate H₂O₂ in millimolar concentrations [33]. Melanocytes are sensitive to reactive oxygen species (ROS) due to low catalase levels in these cells. Kauser et al. [34] verified that catalase expression was reduced both in epidermal and hair follicle melanocytes with age, whereas superoxide dismutases remain constant. The outcome of H₂O₂ accumulation in melanocytes depends on its concentration [35, 36]. Millimolar concentrations of H₂O₂ can damage many proteins and peptides leading to deactivation/disruption of many important pathways involved in melanogenesis. However, tyrosinase as well as other proteins are up-regulated and activated by low concentrations of H₂O₂ [35]. Consequently, catalase expression/activity is getting increasing attention as a key point for the response of melanocytes to ROS. In human melanocytes in vitro, catalase-specific mRNA, protein and enzymatic activity were directly correlated with total cellular melanin content [37]. Because tamoxifen was described as up-regulating catalase gene transcription and catalase production in aorta of rats after one and two weeks of treatment [38], the influence of Tam and 4HO-Tam on CAT gene mRNA levels of NHEM was determined by real-time PCR (Fig. 6). The levels of CAT mRNA in NHEM were significantly higher upon treatment with 4HO-Tam. Tam and ethanol also induced to some extent the transcription of CAT. A correlation between the up-regulation of CAT transcript levels and intracellular melanin induced by physiological concentrations of the two SERMs tested was verified. Some controversy exists in the literature regarding the putative anti-oxidant role of oestrogens and tamoxifen. Tam promotes superoxide production in platelets increasing the thrombosis risk factor [39], and others describe Tam as being able to reduce the Mn-induced ROS formation in astrocytes [28]. The mechanism must be dependent on the cell type and context; however, if SERMs induce ROS production, they can, depending on the ROS concentrations, activate cellular defensive mechanisms or induce oxidative stress. Under the conditions tested, the 4HO-Tam and Tam at physiological concentrations induced catalase expression in NHEM pointing towards a promelanogenic effect mediated by ROS.

Concluding remarks

Based on current knowledge on the beneficial effects of topical application of SERMs on skin ageing, there are still many open questions concerning safety, secondary effects and mechanisms of action. The work here reported demonstrated that Tam and its potent metabolite, 4HO-Tam, have the capacity to induce melanin production and extrusion in normal human melanocytes in vitro, not showing cytotoxicity at physiological concentrations. Melanin itself is a defensive mechanism from photodamage, but uneven pigmentation is undesirable; therefore, further studies must be performed to exclude this hypothesis for topical application of SERMs. The complexity of melanogenesis regulation, which is evident in this work along with other published reports, makes this biological process very hard to modulate exogenously. The mechanism by which these two SERMs act involves changes at the transcript levels of important proteins in the early maturation of melanosomes, PMEL17 and MART-1, and in a newly recognized key melanogene-
sis modulator – catalase. The capacity of inducing anti-oxidant defensive mechanisms is clearly an advantage of these SERMs from the point of view of an anti-ageing cosmetic use. Our results suggest a possible application of SERMs as non-toxic, anti-oxidant and melanogenic compounds to be used in topical formulations in cosmetic industry.

Acknowledgements

Teresa Matamá holds a grant from FCT – Fundação para a Ciência e a Tecnologia (SFRH/BPD/47555/2008). This work was supported by FEDER through POFC – COMPETE and by national funds from FCT through the project PEst-C/BlA/UI4050/2011.

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