Abstract. Skin cancer is one of the most common malignancies affecting humans worldwide, and its incidence is rapidly increasing. The study of skin carcinogenesis is of major interest for both scientific research and clinical practice and the use of in vivo systems may facilitate the investigation of early alterations in the skin and of the mechanisms involved, and may also lead to the development of novel therapeutic strategies for skin cancer. This review outlines several aspects regarding the skin toxicity testing domain in mouse models of chemically induced skin carcinogenesis. There are important strain differences in view of the histological type, development and clinical evolution of the skin tumor, differences reported decades ago and confirmed by our hands-on experience. Using mouse models in preclinical testing is important due to the fact that, at the molecular level, common mechanisms with human cutaneous tumorigenesis are depicted. These animal models resemble human skin cancer development, in that genetic changes caused by carcinogens and pro-inflammatory cytokines, and simultaneous inflammation sustained by pro-inflammatory cytokines and chemokines favor tumor progression. Drugs and environmental conditions can be tested using these animal models, keeping in mind the differences between human and rodent skin physiology.

Contents
1. Introduction
2. Human skin versus animal model skin: Similarities and peculiarities in carcinogenesis
3. Inflammation triggering carcinogenesis
4. History of experimental skin carcinogenesis
5. Refining the model of carcinogenesis
6. Advancing knowledge using models of skin carcinogenesis
7. Testing avenues using animal models of skin carcinogenesis
8. Conclusion and perspectives

1. Introduction

During evolution, the skin has been constantly exposed to complex agents, such as physical (sun rays) and biological assaults (microbes and or allergens); in response to these assaults, the skin has developed specific networks of molecular and cellular sensors to counteract all these aggressors (1). The skin is the largest organ with immune function, and represents an intrinsic network of non-immune and immune cell and molecules that interrelate (2,3). Throughout ontogeny, apart from being exposed to biological aggressors, skin is also exposed to potentially harmful environmental compounds (cigarette smoke, automobile emissions, industrial soot and groundwater) (4).

Due to these many and complex aggressions, skin cancer appears in many forms and it is one of the most common malignancies affecting humans. Owing to various factors, the incidence of skin cancer has increased over the years (5,6). Skin cancer is a multifactorial disease in that it has a strong genetic component. In addition, several environmental factors also play a role in this increased incidence. Among the environmental factors, the most significant is the exposure to ultraviolet (UV) radiation (7), doubled by the exposure to certain chemicals, medication use or stress. All these connected factors may modulate many skin physiopathological processes (8-11), and trigger the initiation and progression of skin cancer (5).
Skin carcinogenesis is a complex, multistep process (12,13), and the study of early alterations in the skin and of the mechanisms involved, as well as the development of novel therapeutic strategies are of interest for both scientific research and clinical practice. Moreover, the striking increase in both the number of new topical chemical entities and their biological effects creates the need to understand the intimate molecular mechanisms that underlie their in vivo effects. Developing in vivo systems for the rapid evaluation of potential drugs is always the concern of researchers. Mouse models of skin carcinogenesis remain one of the most commonly available and cost-effective animal models. In this type of model, agents, either applied topically or systemically, can be studied at the earliest possible stage in the development of drugs/new therapies. In contradiction to the underdosing condition of this model, there is a complex array of mouse strains with distinctive biological behavior to this standard protocol, and this review intends to highlight this distinction and to flag the reported particularities.

2. Human skin versus animal model skin: Similarities and peculiarities in carcinogenesis

Immunologically, mice are the first option when studying immunotoxicology. From this point of view, >15 years ago, the sequencing of human and mice genomes revealed that approximately 300 genes are unique to mice in comparison to humans (14).

A comprehensive review, published >10 years ago, demonstrated that although there are differences in the skin of mice versus human skin, mice are reliable in vivo models that can be used to furnish relevant toxicological information (15). Human therapies are becoming more complex, targeting various cells/proteins/genes; hence, the information gathered from animal models should be carefully weighted in terms of extrapolating data from mice to humans. This is the reason that we have witnessed so many examples of therapies that have had perfect outcomes in experimental animal models, but have then failed to be as effective in humans (16-21).

The main immunological differences regarding the skin is that in mouse skin and mucosa, the predominant T cells are γ/δ T cell receptors (TCRs), whereas in humans, α/β TCRs prevail (22).

T cells that are specific for mouse skin have their TCRs encoded by a single Vγ and Vδ gene. These cells seem oligoclonal (Vγδ Vδ1 genes are encoding for this specific TCR), are found in the epidermis, and they mainly appear as dendritic epidermal T cells. In human skin, T cells with α/β TCRs predominate in human skin, mainly in the dermis. Up to now, TCR γ/δ T cells were not identified in normal human skin (22) and were reported only in lymphoma cases (23). Even in mice, TCR γ/δ phenotype T cells can have different densities between different skin sites and different strains (24). In mice, TCR γ/δ T cells play important roles in tissue homeostasis and during tissue repair. These cells secrete growth factors (e.g., for keratinocytes and insulin-like growth factors), through which the crosstalk between γ/δ T cells from the skin and keratinocytes takes place, actively contributing to the physiology of normal skin and further contributing to wound healing (25). In humans, α/β TCR T cells are activated by CD1 antigen-presenting molecules. It has been demonstrated that a large number of T cells with CD1a-autoreactive phenotype are homed to the skin, producing interleukin (IL)-22 in response to CD1a expressed on a different skin cell population, namely Langerhans cells (LCs) (26). Thus, while in mouse skin T cells collaborate mainly with keratinocytes, in human skin, T cells associate in functionality with specific dendritic cells, such as LCs.

3. Inflammation triggering carcinogenesis

Inflammation involves the secretion of a number of mediators from immune and non-immune cells and occurs for the purpose of the restoration of damaged tissue. Inflammation was known from the beginning of the last century (27), and is a multi-stage process, beginning with the injury inflicted upon a tissue and ending with the reconstruction of the damaged tissue.

Environmental factors can inflict this injury, whether a macro-physical trauma or a micro-trauma (e.g., overuse, friction and sunburn). At the cellular level, the disruption of the cellular membrane releases the intracellular contents into extracellular spaces. Metabolically, hypoxic changes occur, and the cells become deprived of oxygen (secondary hypoxic injury), the sodium pump fails, and cellular membrane disruption continues in adjacent cells, thus enhancing the destruction at a cellular level. This disruption generates mediators (e.g., histamine and bradykinin) that represent the first signals triggering an inflammatory response. The inflammatory response triggers hemodynamic changes: arteries dilate enhancing blood flow, inactive capillaries and venules open, the total blood flow increases, the rate of flow decreases and leukocytes bening to adhere to the vessel wall. Due to critical permeability alterations, leukocytes transmigrate to the injured site. Following the increase in chemoattractant gradient concentration, leukocytes migrate to the injured site. Neutrophils are the first cells that arrive, being the temporary first line of defense [short-lived cells (approximately 7 h)]; neutrophils are followed by macrophages that build up the second line of defense and can survive for months. These two type of cells process, through phagocytosis, debris/microbes, enhancing the clearance process through lymph vessels (28).

In the framework of this review, we focus on a key issue in the process of inflammation, the acute versus the chronic stages of this process (Fig. 1) that can trigger skin carcinogenesis. There is a scientific consensus to incriminate chronic inflammation as linked to carcinogenesis (29) and in mouse experimental models, this chronic inflammation is experimentally sustained in order to favor tumorigenesis.

The differences between these two stages of inflammation reside on different immunological mechanisms. During the acute inflammatory stage, innate cells secrete mediators that attract Th1-type T lymphocytes. Th1 cells secrete cytokines with antitumor activity [e.g., IL-2 and interferon (IFN)-γ]. Apart from T cells, factors secreted by B cells (e.g., antitumor immunoglobulins) activate cytotoxic T lymphocytes (CTLs). Hence, a cellular armamentarium is built up to protect against tumor development.

During the chronic stages of inflammation, when there is a constant activation of the immune response without actually resolving the damage inflicted to tissue, there is an accumulation of regulatory T cells, Th2 cells and activated B cells. In chronic inflammation, immune cells secrete pro-tumorigenic
factors [e.g., IL-4, -6, -10 and -13, and transforming growth factor β (TGF-β)] that inactivate CTL cytotoxicity, thus favoring tumor development (30).

The soluble mediators and cellular effectors of inflammation are common to the tumor microenvironment and can reside in the tumor site. Inflammatory conditions can preclude a malignant transformation and/or an oncogenic alteration sustains the inflammatory microenvironment favorable for tumor development (31), a phenomenon that is exploited in chemically induced skin carcinogenesis.

**Chronic inflammation and tumorogenesis.** In skin there is an established link between tissue damage, inflammation and cancer development. Tumorigenesis is based on constitutive pathway activation, while inflammation is a self-limiting process in normal conditions (32). In the framework of this review, chronic inflammation of the skin can be one of the traits for tumor initiation and progression. In animal models, chronic inflammation may be used to trigger tumorigenesis. The long-term production and the consequent accumulation of inflammatory factors (cytokines/chemokines) can induce locally an immunosuppressive milieu associated with tumor development and progression (29).

Cytokines, long-time players in inflammation, are produced in the skin by an enhanced group of resident cells [keratinocytes, LCs, melanocytes, mast cells (MCs) and macrophages], as well as by recruited inflammatory cells (e.g., neutrophils, eosinophils and lymphocytes) (2). Generally, cytokines are synthesized following cell activation and act locally within the tissue, having a paracrine function in neighboring cells that express specific receptors or have an autocrine action in the cells producing them in an auto-regulatory loop. When the inflammatory stimulus is prolonged, cytokine production becomes excessive, and finally, this has a deleterious effect both on site and on distal sites from the original inflammation site, similar to the effects of hormones. Cytokine receptors have a mainly homologous structure; thus, various cytokines can have pleiotropic effects, acting on several targets. Beyond that, cytokines can have a synergistic effect on the same cells, while acting antagonistically on other cell types (33).

As it has already been acknowledged, in each tissue, as with the skin, a complex cytokine network is developed and this network is precisely regulated (33,34); any deregulation of this cascade can trigger a tumorigenic process (Fig. 2).

Cell migration is critical for several normal processes, such as embryogenesis, the immune response, inflammation; however, it is also one of the key events in cancer cell metastasis (35). All the molecular perturbations underlying chronic inflammation as triggers for skin tumorogenesis are again the center of cell migration research. Ehm2 (a novel cancer promoter) belongs...
to the FERM family of proteins (4.1 protein, ezrin, radixin, moesin), that are involved in membrane-cytoskeletal interactions, and are linked to the metastatic event in several types of cancer, including skin cancer. In a study published in 2013, the effect of Ehm2 knockdown on cell migration, adhesion, growth, cell cycle progression and apoptosis were reported. The authors demonstrated that Ehm2 was expressed at 3-fold higher levels in tissues with acute inflammation, compared to the chronic state. Following the knockdown of Ehm2, the expression of another protein was found to be decreased, namely that of neural Wiskott-Aldrich syndrome protein (Nwasp) (36). Nwasp is significant in the invasion processes as the binding of cortactin to Arp2/3 and Nwasp are key elements for invadopodia formation in melanoma cells (37). These reported results propose that Ehm2 proteins are interesting connection molecules between inflammation and the skin cancer metastatic process. In melanoma, following the knockdown of Ehm2 protein, the expression of Nwasp was downregulated, directly affecting cell migration (36).

There are intracellular pathways that are deregulated in tumorigenesis and in a non-healing wound. Thus, both processes involve common molecular and signaling pathways, such as Ras, Hedgehog and WNT (38).

Epithelial-mesenchymal transition (EMT) is a process through which epithelial cells lose some of their characteristics (cell polarity and cell-cell adhesion) and gain migratory and invasive properties, becoming mesenchymal stem cells (39). When a wound is healing, the EMT process is activated (40). Epidermal keratinocytes acquire migratory phenotypes (41) and then return to normal upon wound closure during the rebuilding of the basement membrane. In tumors, epithelial cells can harbor oncogenic mutations, undergoing EMT, a process associated with the acquisition of cancer stem cell properties (42). Another recently discovered cellular process, transdifferentiation, is a mechanism that governs the transformation of a mature somatic cell in another mature somatic cell lacking the intermediate pluripotent state or progenitor cell type phases (43). In adult tissues, transdifferentiation is not as accelerated as in more immature ones; however, in wounding, mesenchymal stem cells undergo transdifferentiation into epidermal cells, endothelial cells and pericytes (44,45).

Another mechanism that can link inflammation with tumorigenesis is the fact that during wound healing, fibroblasts deposit excess collagen (during fibrosis), which leads to scar formation. This connective tissue is a microenvironment that is tumor permissive (46). As stated above, in this microenvironment, the presence/production of growth factors, cytokines and chemokines is similar in chronic wounds as in tumors, with a slight difference being in the expression levels in terms of kinetics (31) (see also Fig. 2).
In conclusion, we can ascertain that a close association exists between chronic tissue damage, inflammation and cancer (32); in this context tumors may develop, although uncommonly, also at the site of chronic skin wounds (47).

4. History of experimental skin carcinogenesis

Where the story begins. One of the first recorded studies regarding two-step chemically induced carcinogenesis is that of Frei and Stephens published >45 years ago (48). It was actually the first study to demonstrate that tumors induced in mouse ears correlate with the induction of hyperplasia as an inflammatory response. Leucocytes migrating to the site were found to correlate with the rate of induction of hyperplasia; however, at the same time, no clear association between inflammation, hyperplasia and tumorigenesis was observed.

From these decisive results, chemically induced mouse skin carcinogenesis was the main animal model of cutaneous tumorigenesis (49‑51). This model was used for evaluating anti-tumor drugs, but also for understanding the nature of epithelial cancers as a multi-stage process (52).

Several protocols have been developed for ‘two-stage’ carcinogenesis in which tumor incidence, tumoral latency, multi-staging and the progression of the skin cancer are studied. In the model of two-stage skin carcinogenesis, the tumor is initiated after a single sub-carcinogenic dose of 7,12-dimethylbenz[a]-anthracene (DMBA). This irreversible event leads to visible tumors only after the repeated application of a promoter agent, such as the phorbol ester, 12-O-tetradecanoylphorbol-13-acetate (TPA). Therefore, unlike one-step carcinogenesis, in two-stage carcinogenesis, the initiation and promotion phases can be noticeably separated, both functionally and temporally (52). This distinction of phases offers a tremendous advantage when studying the effects of environmental factors and/or drugs in the different stages of tumorigenesis.

Another advantage offered by this two-stage model is that tumor development can be visually monitored during the life-span of a mouse. With this dynamic monitoring, non-invasive evaluation methods can be used (see below for images of in vivo confocal microscopy). Moreover, at the end of the experimental period, the transformed tissue can be harvested and thoroughly examined. This animal model yields good reproducible results; thus, when assessing cytokinetics, chemopreventive and/or dietary agents for skin tumorigenesis, reliable results are obtained (53).

Over the past years, genes and cell signaling pathways, the molecular mechanisms that underlie tumorigenesis, were studied using this animal model (54,55).

Considering the broad utilization of this animal model, its flexibility to extensive experimental approaches, as well as its unproblematic development for any animal husbandry, two-stage skin carcinogenesis developed in mice can be used for the model of human carcinogenesis when cancers of epithelial origin are studied (56).

Strain differences. One of the first studies acknowledging different strain susceptibility was published >30 years ago (57). In this early study, authors used two-stage skin carcinogenesis with a polycyclic aromatic hydrocarbon (DMBA) as the initiating carcinogen, and subsequently, as the promoter of carcinogenesis, croton oil. The study used LACA and BALB/c mice, namely one susceptible and one resistant strain. In this early study, the authors reported that DMBA was more effective in the LACA strain compared to the BALB/c one. Moreover, this was one of the first studies demonstrating the metabolic activation of DMBA and its transformation in the active carcinogen, a process common to both mouse strains, whether resistant or sensitive. This early study flagged strain differences in terms of the DMBA response, although DMBA was the actual promoter of carcinogenesis in both studied strains. At that time, mechanisms such as DNA repair were still under investigation and thus, strain differences as regards the induction of carcinogenesis could be only suggested to involve these mechanisms. Dissimilar to carcinogens, such as DMBA, phorbol ester promoters (component of the utilized croton oil) appear to shunt any metabolic activation, while being degraded and inactivated by epidermal cells. These early experiments with different promoters did not provide any evidence that the distinct biological behavior of different mouse strains is sustained by different promoter degradation (57).

After another 10 years, the era of transgenic mice began and the subject of strain specificity in terms of developing chemically induced carcinogenesis gained another dimension. The role of specific genes as determinants for particular behavior was acknowledged. Strain differences in terms of susceptibility to specific toxic agents are in fact an in vivo tool to be exploited and, in this light, transgenic animals that comprise specific genes can be manipulated. At that time, several experimental pathways were highlighted, pathways that now are at the post-graduate research level. It was then stated that transgenic animals can be used in several ways: to introduce a human gene encoding a drug metabolizing enzyme, or to delete and/or modulate the expression patterns of specific within target cells (58).

In these early experiments, it was shown that the promoter of a stress-regulated gene linked to a reporter gene (lacZ or green fluorescent protein) can be inserted into the mouse genome; then upon experimental tumorigenesis, sensitive cells can be identified as the cells are carrying the reporter gene. Transgenic mice can be used for the high-throughput screening of compounds, an invaluable experimental model that can offer results regarding the tissue and cellular specificity of certain drugs (58). As early as the 1990s, authors highlighted that transgenic animal studies are complex, and that altering one gene does not mean one straightforward protein alteration. The introduction of a specific human cytochrome P450 gene can have no metabolic effect due to the overall background activity in the rodent. Drug uptake, metabolism, detoxification and repair, as parts of the complex process of toxicity, differ in primates and rodents; for example, a toxic response in one species can be totally irrelevant in another (58); however, transgenic animals in the toxicological domain have gained momentum and are speeding up the drug discovery processes.

Almost at the same time period, another team investigated the association between single gene mutations and carcinogenesis, the histological type triggered by this gene mutation and whether a novel papillomavirus (at that time) had a co-carcinogenic function (59). A two-step protocol (DMBA followed by TPA) was used to induce papillomas in several
inbred, hybrid, and various genetically altered mice. Studying the histological types of tumors, the authors reported an increased panel of histology beginning with early follicular papillomas, along with mixed papillomas, or exophytic papillomas, hyperplastic papillomas, fibropapillomas, squamous cell carcinomas (SCCs), and MC tumors. Moreover, in these mouse models, high-, intermediate-, and non-responding groups can be obtained. The authors demonstrated that tumor induction was conditioned by skin-specific mutant genes. Papillomavirus antigens or viral genomic DNA was not present in the induced tumors. Genetic differences were first flagged in the late 1990s as underlying the strain differences (59).

Laboratory animals subjected to specific genetic manipulations can be used to not only study compounds used to induce carcinogenesis, but also to study the effects of topical chemical toxins. This was an important conclusion of the late 1990s, paving the way for the future genomic era of skin toxicology and dermato-oncology.

5. Refining the model of carcinogenesis

As discussed in the previous section, several decades of studies have investigated the induction of carcinogenesis in animal models. The topical administration of compounds which induce skin carcinogenesis in mice provides an easy model for studying local, systemic and environmental factors influencing tumor susceptibility, growth and progression. This model of chemically induced carcinogenesis almost as old as modern dermato-oncology and skin toxicology continues to provide a cost-effective model needed for the identification of biological, immunological and molecular pathways implicated in skin tumorigenesis. Besides these somewhat fundamental findings, this model is frequently used for topical antitumoral drug testing. For instance, the two-stage mouse carcinogenesis model was typically used for testing drugs effective for skin cancer prevention (60,61). Studies regarding chemoprevention testing were reported in the SENCAR mouse, a model evaluating phenylretinamides as tumor suppressors via retinoid receptor-independent mechanism(s) (62).

In essence, this model uses the two-stage application of chemicals to the skin for the initiation and promotion of cutaneous tumors. When using a two-stage model of induction, after a single application of the initiator mutagen, DMBA, repeated applications of a pro-inflammatory phorbol ester (TPA), or phorbol 12-myristate 13-acetate (PMA) are carried out. The literature indicates that tumors that appear can be benign papillomas that regress or progress to SCC (63). Moreover, the direct appearance of SCC was reported without the presence of any pre-cancerous lesions. Thus, two-stage chemically induced carcinogenesis provides an opportunity to monitor early and late events in cancer development and progression (63).

Our experience has shown that, when we used this protocol (64), in three different mouse strains, widely covering the susceptibility area in terms of chemically induced carcinogenesis (high susceptibility, nude mouse CD1-Foxn 1nu strain; medium susceptibility, CD-1 strain; and low susceptibility, C57BL/6 strain) we obtained various results. Thus, from the C57BL/6 mice, only a small percentage (5%) developed skin lesions. Tumor formation began approximately after 25 weeks from the first DMBA application. Macroscopically, they displayed only one formation, round shaped, rough at palpation with a wide base. We did not observe metastases at necropsy and the histopathological identification (Fig. 3) revealed a poorly differentiated carcinoma. In order to maintain the homogeneity of the mouse model, we established a clone of tumor cells with which we established an in vivo mouse model of skin carcinoma using the C57BL/6 mouse strain.

In the CD1-Foxn 1nu mouse group, approximately 20% of the mice survived during the experiment. In the survivors, tumor formations appeared around 8 weeks from the first DMBA application and the mice were sacrificed 10 weeks after the appearance of the lesions. Macroscopically, the mice developed multi-formations, 1-3 mm in diameter, with an irregular appearance and a narrow base. No metastases were observed at necropsy and the histological examination depicted cutaneous papillomas (Fig. 4). All the presented original animal experiments were carried out in accordance with EU guidelines (65).

The mice in the CD1 group all developed skin formations. In this case, the tumor was clinically detectable as early as 5-6 weeks after the initial application. Macroscopically, there were multi formations, elongated with an irregular shape, soft at palpation, with a narrow base and a rapid growth rate. No metastases were observed at necropsy. A technology that we and others are developing in animal models is in vivo reflectance confocal microscopy (RCM). This modern imaging technique allows for the non-invasive ‘quasi-histological’, real-time
endogenously carcinogenesis and that TNF-α promotes tumor necrosis factor-α-inducing carcinogenesis points toward the role of TNF-α in tumor-promoting inflammation, this updated study involves another factor in carcinogenesis which is chemically induced, but concomitantly shows that these characteristics are common to humans as well. In making a comparison of the differences in the skin tumor microenvironment between the mouse model and humans, common molecular mechanisms were depicted and the authors recommend the use of mouse models in skin toxicological testing (75). In C3H/HeN mice, the exposure of the skin to UV radiation induced the activation of β-catenin in a time- and dose-dependent manner. Mice deficient in cyclooxygenase-2 (COX-2) subjected to UV radiation exhibited a lesser activation of β-catenin. In SKH-1 hairless mice exposed to UV radiation, the activation of β-catenin signaling was also observed as well as the development of UV radiation-induced skin tumors. All the accumulated data from the study of photo-carcinogenesis led to the conclusion that the activation of the β-catenin pathway can induce skin carcinogenesis (76).

6. Advancing knowledge using models of skin carcinogenesis

Genomics. The majority of publications in the genomics investigating field have focused on studying the environmental factors, such as UV radiation and carcinogens as potent inducers of epigenetic alterations. Exposure to environmental mutagens or simply spontaneous errors can induce somatic mutations, leading to the development and/or progression of cancer. The epigenome can also undergo chemically induced or spontaneous alterations, leading towards carcinogenesis. Next-generation sequencing can identify genetic variants, somatic mutations, gene expression profiles, and epigenetic alterations with single-base resolution. All this new technology was put in use to provide insight into tumorigenesis driven by environmental factors (77).

Moreover, nutritional and dietary resources can modify individual susceptibility through changes in the epigenome. Epigenetics hence aims to identify the health risks of environmental toxicants combined with dietary and nutritional ones. The epigenetic end-points reported by authors highlight global hypomethylation and specific hypermethylation at diverse tumor suppressor genes upon environmental exposure. Furthermore, a series of conditions were flagged that could influence the epigenetic effects of environmental factors: namely the dynamics of exposure, dose, gender, organ-target and age were examined for alterations in the epigenome. The effects of environmental factors can be balanced by nutritional/dietary agents, reversing their epimutagenic effects (77).

In a study published in 2014, the genome-wide DNA methylation profile was investigated in a model of UVB- and DMBA/TPA-induced skin cancer (78). In that study, the SKH-1 strain was subjected to UV radiation and in CD-1 mice, carcinogenesis was induced by DMBA/TPA. The authors reported >6,000 genes in the UVB group and >5,400 genes in the DMBA/TPA group that proved an enhanced CpG methylation. Using ingenuity pathway analysis, the first pathways in which these deregulated methylation appeared were the ones frequent HA-RAS-mutated tumors are reported in melanoma following treatment with B-raf inhibitors. This recent finding indicates the probable existence of HA-RAS-mutated cancer stem cells in the actual tumors. In a similar manner, UV-induced human SCC displays mutations in TP53, a tumor suppressor gene. This published review emphasizes the differences in skin tumorigenesis which is chemically induced, but concomitantly shows that these characteristics are common to humans as well. In making a comparison of the differences in the skin tumor microenvironment between the mouse model and humans, common molecular mechanisms were depicted and the authors recommend the use of mouse models in skin toxicological testing (75). In C3H/HeN mice, the exposure of the skin to UV radiation induced the activation of β-catenin in a time- and dose-dependent manner. Mice deficient in cyclooxygenase-2 (COX-2) subjected to UV radiation exhibited a lesser activation of β-catenin. In SKH-1 hairless mice exposed to UV radiation, the activation of β-catenin signaling was also observed as well as the development of UV radiation-induced skin tumors. All the accumulated data from the study of photo-carcinogenesis led to the conclusion that the activation of the β-catenin pathway can induce skin carcinogenesis (76).
related to tumorigenesis. Hence, of the deregulated pathways, cAMP-mediated signaling, G protein-coupled receptor signaling, and PTEN signaling were associated with UV radiation, while protein kinase A (PKA) signaling and xenobiotic metabolism signaling were associated with DMBA/TPA carcinogenesis. As stated in the inflammation-carcinogenesis section, the inflammatory processes can sustain tumorigenesis (78).

Thus, in that study, IL-6-related inflammatory pathways had alterations in the methylation profiles when skin was subjected to UVB irradiation. Altered genes were classified in the UVB alterations in the methylation profiles when skin was subjected to environmental factors and DmBA/TPA models, while establishing in silico their molecular interaction networks. The authors demonstrated that methylation profiles of skin subjected to environmental factors (UVB irradiation or chemical carcinogenesis DMBA/TPA) can shed light on epigenetic gene regulation in skin carcinogenesis (78).

Actually the first study on the aberrant methylation of Nr4a3 exon 3 CpG island was reported in a multistep mouse model of skin carcinogenesis (79).

Studies published in 2014 demonstrated that many cancer susceptibility loci are located throughout the genome (80,81). Mapping these loci in a mouse model of skin cancer, the authors demonstrated strong genetic loci that render resistance to chemically induced skin papillomas. These loci are located on chromosomes 4 and 7. Combining congenic mapping (congenic mouse strain, FVB.MS-STMm3) and allele-specific alteration analysis of these loci located on chromosome 4, it was demonstrated that STMm3 (skin tumor modifier of MS3) responsible genes influence the formation of papillomas in two-stage skin carcinogenesis by regulating papilloma growth rather than development (80). The same group, in the same year, demonstrated that STMm1 may be responsible for papillomagenesis in two-stage skin carcinogenesis by regulating epidermal quiescent stem cells (81).

Genetically modified animals have been used as valuable tools in depicting the carcinogenic process. A number of studies have used these animals in the animal model of two-stage skin carcinogenesis in order to depict the roles of certain genes involved in epithelial carcinogenesis (54,82,83). In these animals, the functionality of genes can be studied throughout the evolution of the carcinogenesis process. Targeting genetic modifications that can modify cutaneous tissue has allowed different genetic manipulations (gene overexpression or gene deletion) in particular skin compartments, recapitulating spontaneous genetic events in tumorigenesis (84).

In the search for new therapeutic targets from the panel of signaling pathway molecules, transgenic mouse have been used. The study by Wilker et al demonstrated that using LY294002 (a PI3 K inhibitor), in a transgenic mouse model that overexpressed human insulin-like growth factor-1 (IGF-1), emphasized the role of PI3 kinase and Akt-mediated signaling in epithelial carcinogenesis (85).

MicroRNAs (miRNAs or miRs) as regulators of gene expression came into the spotlight in the last 10 years (86,87). Therefore, it was observed that the exposure to different carcinogens also alters miRNA expression (88–90). miRNA expression differs in the different stages of chemically induced carcinogenesis. Carcinogenic agents are potent miRNA expression deregulators, while non-carcinogenic chemicals have a lesser influence on miRNAs. Their expression also differs in different biological systems. Hence, there are increased changes in cancer-target tissues in comparison to the non-target tissues following an acute or a chronic exposure to carcinogens. The families of deregulated miRNAs in carcinogens regulate genes involved in xenobiotic metabolism, carcinogen-induced hypomethylation, DNA repair, apoptosis, cell proliferation, tumor suppression, cell transformation, oncogenesis, tumor angiogenesis, tumor progress and malignant transformation (91). Moreover, there are miRNAs with double functions, such as putative oncogenes or tumor suppressor genes (92). Carcinogens can influence the balance between these functions and drive their functions toward tumorigenesis. As recently demonstrated, miRNAs specific to carcinogen exposure can be used as biomarkers for genotoxicity and carcinogenicity (93).

The clear message conveyed from these data is that new endpoints of environmental toxicants should be found and that more attention should be paid to finding measure to prevent environment-related diseases.

Proteomics. As the domain of proteomics is rapidly evolving, mouse models of chemically induced carcinogenesis have also taken advantage of this area. The differences in the susceptibility between strains reside in the genetic diversity that triggers protein differences. In a recent study, specific protein or signaling pathway alterations were investigated in association to this susceptibility (94). Examining the epidermis proteome in DBA/2 sensitive and C57BL/6-resistant mice upon the administration of TPA, 19 differentially expressed proteins were found. Five proteins were calcium-binding proteins: annexin A1, parvalbumin α, S100A8, S100A9 and S100A11. The S100A8 and S100A9 protein levels were found to be increased when using a different mouse model with the topical application of tumor promoters, okadaic acid and chrysoarobin. When examining the association between these 19 upregulated proteins, it was shown that they are active players in several inflammatory networks involved in skin tumor promotion, such as TNF-α and nuclear factor (NF)-κB. Nucleic acid miRNAs for various proteins such as TNF, Nfkβ1, IL-22, IL-1b, and chemokines such as Cxcl1, Cxcl2 and Cxcl5 were found to be highly expressed following the administration of TPA in DBA/2 mice in comparison to C57BL/6 mice. These reported results indicate that inflammatory genes can sustain the basis of genetic differences in susceptibility in mouse models of chemically induced carcinogenesis (94). Apart from inflammatory processes that can account for these different effects, tetraspanins, cell-surface proteins present on almost all cell and tissue types have been shown to be involved (95). During two-stage mouse skin chemical carcinogenesis, CD151 favors tumorigenesis, inducing a more rapid development, multiplicity, size and progression to SCC in this model. In human skin developing SCC, CD151 expression is increased. CD151 is associated with the activation of the transcription factor, signal transducer and activator of transcription 3 (STAT3), and PKCα-α6β4 integrin. CD151-PKCα is associated with a more invasive SCC. In mouse models at least, CD151 was proven to be a future antitumor therapy target (96). In a recent study, in a model of DMBA-induced skin carcinogenesis using PKCα knockout mice, it was shown that PKCα suppresses tumor formation, but not tumor growth and progression to skin carcinogenesis (97). Using IL-6 and TNF-α-deficient mice in chemically induced skin carcinogenesis, it was also demon-
strated that IL-6 is not the main pro-inflammatory cytokine that promotes tumorigenesis of the skin. The authors concluded using this model that individual cytokines have distinct and discrete roles in tumor promotion (98).

Proteomics still has an important input in the nearby future in this domain as its rapid evolving methodologies find their place in quantifying the proteome behind the subtle process of tumorigenesis (99,100).

7. Testing avenues using animal models of skin carcinogenesis

Determining how close we are to the human scenario. The presented animal model runs through the characteristics of human multi-stage carcinogenesis. As reported several years ago, it seems that mutations within stem cell niches can be the initial step in the flow of events leading to tumorigenesis (101,102). At the genetic level, mutations and genes that are activated resemble those identified in humans. Therefore, when mutations in ras family members appear, receptors of the epidermal growth factor (EGFR) are activated, signaling pathways of Stat3 and Akt-mediated are activated and the mutations of TGF-β1 and Tp53 occur, and these are all genetic processes that seem common in both species (49).

As humans are subjected to multiple carcinogenic doses, mainly accumulating low doses, this two-stage skin carcinogenesis protocol uses both carcinogens and promoting agents, as in an environmental scenario. Furthermore, in human cancers, there is a long latency that is also mimicked by the promotional component for tumor development (103). As a result, there is a commonly agreed opinion that this mouse model can be utilized to study the mechanistic basis of human epithelial cancers.

Drug testing using the model of skin carcinogenesis. The toxicological drug testing area is a domain that frequently uses this mouse model.

The effect of nano-sized titanium dioxide particles (TiO₂), as a growing component used in cosmetics, sunscreens and food additives, was tested in the model of two-stage skin chemical carcinogenesis using DMBA and afterwards the tested compound (104). The topical application of non-coated rutile-type TiO₂ did not trigger any UV-induced skin carcinogenesis in rats, probably due to the lack of penetration of TiO₂ into the epidermis. The same authors switched to chemically induced skin carcinogenesis to test the effect of silicone-coated TiO₂. They used mice, human c-Ha-ras proto-oncogene transgenic mice (rasH2) and resistant CD-1 ones. The compound did not influence the DMBA carcinogenesis in sensitive mice or resistant mice, possibly due to the lack of penetration through the epidermis (104).

In another study, the efficacy of topically applied liposome-encapsulated tamoxifen (TAM) in a model of DMBA-TPA-induced skin tumorigenesis in order to potentially decrease its systemic toxicity was tested. TAM was encapsulated in special liposomes and the incidence of papillomas was examined (105). This recent study demonstrated the inhibition of skin carcinogenesis when liposome conditioned drug was used, this finding bringing good news for the future use of liposomal systems/drugs in skin cancer (105).

Agaro-oligosaccharides (AGOs) studied in in vitro models have been proven to suppress nitric oxide (NO) levels, as well as the production of prostaglandin E (PGE) and pro-inflammatory cytokines (106). Continuing this endeavour, in the two-stage mouse model of skin carcinogenesis, the administration of AGOs delayed the appearance of tumors and decreased the number of tumors. PGE2 was shown to be suppressed by AGO intake in a model of TPA-induced ear edema, while COX-2 and microsomal PGE synthase-1 were found to be downregulated. As stated above, this was a mouse model that reported results of the anti-tumor effects of AGOs, as well as of the anti-inflammatory, PGE-mediated effects (106).

As demonstrated in another study, the topical administration of the βG inhibitor, D-glucaro-1,4-lactone (1,4-GL), or D-glucuronic acid-γ-lactone (GUL) its precursor, to SENCAR mice with skin carcinogenesis, epidermal hyperplasia was significantly reduced along with a decrease in inflammation (107). Moreover, Ha-ras mutations were reduced upon experimental therapy. That study demonstrated that whether administered topically or through diet, therapy with GUL or 1,4-GL was effective in hindering experimental skin tumorigenesis (107).

Natural products testing. As there is a continuous effort to establish dietary routes of harmful compounds (108), over the past years, several papers were published showing results of natural compound testing in animal models of skin carcinogenesis and these models are particularly suited for the evaluation of the effects of dietary factors/dietary manipulations on tumor development.

Potential natural compounds can be evaluated for their effects on cutaneous tumor initiation, promotion and/or progression; several years ago, a study was published on the anti-inflammatory effects of resveratrol in this model (109). Kleiner et al also showed that the delivery of isopimpinellin from citrus, prior to exposure to DMBA, significantly inhibited tumor initiation (110). Singh et al showed that silymarin was effective at blocking tumor formation and at promoting even the regression of established tumors, this effect being dependent on the sequence of delivery (111). The topical application of rapamycin was reported as a chemopreventive agent, leading to the regression of papillomas in a model of chemically induced carcinogenesis (55,112).

For testing the hydroalcoholic extract of Brazilian red propolis (HERP), an animal model of of dermal carcinogenesis was used. A single DMBA agent was used and animals treated with HERP exhibited significantly decreased tumor multiplicity throughout the 5 weeks of tumor promotion. Administered orally in a murine model of chemically induced SCC, this natural product exerted a significant modulatory effect on the formation, differentiation and progression of tumors (113).

In a previous study (114), geraniol, an acyclic monoterpene alcohol, was tested in a mouse model of DMBA-induced skin carcinogenesis without any inflammatory compound, and skin tumors occurred in all mice. The animals were orally administered geraniol and monitored for lipid peroxidation and antioxidant status. The yielded revealed positive results; geraniol at certain doses prevented tumor formation and regulated the antioxidant status. As the authors explained, the model yielded clear-cut results; geraniol inhibited tumor cell
proliferation, modulated detoxification agents and enhanced free radical scavenging (114).

In another study, in Swiss albino mice exposed to DMBA, the chemopreventive efficacy of rosmarinic acid was reported (115). As in the above-mentioned study, phase I and II detoxification agents, lipid peroxidation by-products, antioxidants and apoptotic biomarkers were assessed. The authors reported that they succeeded in inducing SCC in all the mice within 15 weeks of topically applying only DMBA. The investigated parameters were found modified in the SCC tumor-bearing animals, while extremely positive clinical results were obtained in the animals treated orally with rosmarinic acid. The treated animals did not develop tumors, while animals already bearing SCC tumors exhibited a normalization of all the tested detoxification agents, lipid peroxidation by-products, antioxidants and apoptotic markers (p53, Bcl-2, caspase-3 and caspase-9) (115).

SHR mice subjected to DMBA protocol were also tested for the efficacy of melatonin and metformin. The published results indicated that melatonin and metformin, significantly attenuated tumorigenesis and decreased the overall lipid peroxidation levels (116).

Green tea is also a natural product that exerts beneficial effects on skin carcinogenesis in animal models. Green tea polyphenols (GTPs), mainly epigallocatechin-3-gallate (EGCG) were tested, among other types of cancer, in skin cancer models as well. It was demonstrated that GTPs/EGCG can promote anti-neoplastic processes, apoptosis, cell cycle arrest and can suppress metastasis in tumor cells, but not in normal cells. The reported difference possibly reside in the molecular mechanisms triggered by GTPs/EGCG in signaling pathways in transformed cells versus normal ones (117).

This model is also suitable for investigating the effects of dietary intake on carcinogenesis as calorific restriction is known to inhibit/delay the phase in which the tumor is promoted (118). The altered energy balance that can induce tissue-specific alterations, along with systemic effects can be evaluated in this type of animal model (119,120).

Studies on the chemoprevention of skin cancer have taken advantage of the two-step mouse model of carcinogenesis. The plant flavonoid, silymarin, was proven to be effective against chemical/photo carcinogenesis. In several models of skin carcinogenesis, the topical administration of silymarin inhibited the effects of DMBA, as well as those of tumor promoters, such as TPA, mezerein, benzoyle peroxide and okadaic acid (121). The same chemopreventive effects were obtained for UVB-induced skin carcinogenesis (122). It seems that the actual effects of this flavonoid are its antioxidant, anti-inflammatory and immunomodulatory characteristics that hinder inflammation and evolution to tumorigenesis. In clinical studies, this flavonoid was reported to reduce the toxicity associated with chemotherapy (123).

Taking into account the data from recent reports, it can be emphasized that in the last couple of years, the study of the topical and systemic antitumor effects of ‘green’ drugs has benefitted from this mouse model. As in any study on chemoprevention or dietary intervention, the efficacy of the test compound/drug studied should be carefully examined and coherent conclusions should be drawn for proper translation into human pathology.

8. Conclusion and perspectives

We can outline several aspects in the dermato-oncology and skin toxicology testing area that uses chemically induced skin carcinogenesis. Thus, there are both pros and cons for the utilization of this model.

There are important strain differences in histological type, development and clinical evolution of the skin tumor. These differences are enhanced by the fact that human skin tumorigenesis can comprise other pathways and we should be aware of the existing limitations. In this methodology, mice develop primarily papillomas, without any direct human equivalent. The subsequent SCC tumors that develop through tumorigenesis histologically resemble human SCC tumors. For the chemical initiation, in mice, Hras is the primary target, while in humans, Tp53 appears to bear the gene mutation in human non-melanoma skin cancer (124). The Hras gene from mouse skin closely resembles the gene found mutated in other human epithelial cancers (lung, colon and pancreatic cancers) (125). The last draw-back is that the two-stage model of skin carcinogenesis model is limited when studying metastasis, as the rate of metastatic cutaneous tumors is quite low (126).

The advantages of this model reside in the fact that upon genetic manipulation, the model can mimic the exact genetic and phenotypic changes of certain skin disease conditions. By doing so, drug target validation, preclinical testing and biomarker discovery can take advantage of rapid and oriented results. The roles of potential cancer risk modifier genes, such as proto-oncogenes, and/or tumor suppressor genes in complex processes such as initiation, promotion and progression can be additionally studied in these murine models.

Despite the stated differences, authors recommend the using of mouse models for skin toxicology testing because at the molecular level, common mechanisms are depicted. As in human cancer development, genetic alterations caused by carcinogens and pro-inflammatory cytokines, simultaneous inflammation sustained by pro-inflammatory cytokines and chemokines favor tumor progression.

One of the main issues in testing environmental factors and/or drugs is the similarities between the effects observed in vitro versus the ones registered in vivo. Moreover, another outstanding issue is to what extent we can rely on the experimental animal models when extrapolating the results to humans.

The newly released ToxCast phase II results, demonstrate that high-throughput assays for characterizing rodent toxicants, can integrate both bioactivity and chemical structure. ToxCast phase I has shown that research on drugs combined with the intensive use of omics technologies, are mandatory for combining biological and chemical information in exploring the in vitro to in vivo connection (127).

This domain still needs to explore in depth the interrelation between cells of the immune system and tumor cells, in the continuous search to unravel the intimate molecular mechanisms that trigger skin tumorigenesis (128,129).

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