CORRELATIONS BETWEEN PRECANCEROUS LESIONS AND MELANOCYTIC AND NON-MELANOCYTIC SKIN CANCERS

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Abstract: Nonmelanocytic skin cancers (NMSC) are the most common frequent of skin cancers, which may appear de novo, in normal skin or on precancerous cutaneous lesions. There are many pre-existing lesions holding a potential to transform in skin cancers. Of these, the most common are actinic and seborrheic keratoses, sebaceous nevus, leukoplakia etc. Malignant melanoma (MM) can appear after a malignant transformation of melanocytic lesions or of nevi with a dysplastic potential or de novo in apparently healthy skin. Lately, the medical community has observed more frequent de novo MM than the ones building upon underlying nevi lesions. Based on these data and taking into account that the diagnosis and treatment of precancerous lesions would allow reducing the incidence of skin malignancies and thus mortality, especially in MSC, we conducted a prospective study in which we analysed precancerous lesions and the MM and MSC that appeared on these.

INTRODUCTION

Malignant transformation of precancerous lesions is suspected when there are changes in the shape, size, colour and texture of the lesions. In the case of NMSC, the malignant transformation of precancerous lesions takes a long period of time, sometimes even decades, unlike the MSC, where changes can occur very rapidly.

Precancerous lesions that are most commonly transformed in basal cell carcinomas (BCC) and squamous cell carcinoma (SCC) are actinic keratoses, seborrhic rarely, leukoplakia (specifically for SCC lip) etc. In 2009, Criscione et al. published a study on a total of 7784 of actinic keratoses, observing that the risk of transformation in SCC ranged from 0.6% after 1 year and 2.57% after four years, and that the risk of transformation in BCC varied between 0.48% after 1 year and 1.97% after 4 years.1

Where MM appeared on precancerous lesions, it was observed that the lesions were atypical nevi structures, in which, at a certain moment, architectural and structural changes occurred, drifting towards malignancy. In existing studies, over 50% of MM appears on existing nevi lesions, the percentage ranging and depending on clinical or histopathological criteria. By corroborating clinical and histopathological criteria, it is considered that 39.5% of MM appears upon pre-existing structures of nevi.2 The importance of the diagnosis and treatment of precancerous lesions lies in the possibility of decreasing the incidence and mortality of skin cancer. Since NMSC are the most common of skin cancers, the decrease in their incidence will allow an earlier diagnosis, treatment in the early stages and a total reduction of costs, which could alternatively be used for the initiation of dermato-oncologic education programmes.

METHODS

We conducted a prospective study over a period of three years in cases of MSC and NMSC in the department and outpatient of Dermatology and Oncology from the Clinical Hospital, Sibiu. 385 patients with skin tumours were kept under observation, including 275 with BCC, 57 SCC and 83 with MM. We made a questionnaire for patients affected by NMSC and MSC - the data were extracted from the observation sheets and the dermato-oncologic sheets of patients with skin tumours.

In our group of patients, in addition to the demographic data, we followed the emergence of MSC and NMSC upon new or preexisting lesions, as well as the correlation between precancerous lesions and the type of skin tumour. Data were statistically analyzed using SPSS version 10.

RESULTS

Of the total of 385 skin tumours included in our study most were NMSC (78.44%), while MSC constituted the rest (21.56%). The most common skin tumours were BCC, representing 81.12% of NMSC and 63.64% of patients from group 1; SCC tumours were at 18.87% of NMSC and 14.80% of

Keywords: precancerous lesions, nonmelanoma skin cancers, melanocytic skin cancers

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the total skin tumours (figure no. 1).

Figure no. 1. Group distribution upon the type of tumour

![Diagram showing group distribution of tumour types.](image)

### Table no. 1. Analysis of precancerous lesions in the group of patients

<table>
<thead>
<tr>
<th>Preexisting lesions</th>
<th>BCC</th>
<th>SCC</th>
<th>MM</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>De novo</td>
<td>207 (84.49%)</td>
<td>46 (80.70%)</td>
<td>40 (59.04%)</td>
<td>302 (78.44%)</td>
</tr>
<tr>
<td>Actinic keratoses</td>
<td>18 (7.35%)</td>
<td>4 (7.03%)</td>
<td>0</td>
<td>22 (6.38%)</td>
</tr>
<tr>
<td>Scars</td>
<td>20 (8.16%)</td>
<td>1 (1.75%)</td>
<td>0</td>
<td>21 (5.45%)</td>
</tr>
<tr>
<td>Cheilitis</td>
<td>0</td>
<td>5 (8.77%)</td>
<td>0</td>
<td>5 (1.30%)</td>
</tr>
<tr>
<td>Ulcers</td>
<td>0</td>
<td>1 (1.75%)</td>
<td>0</td>
<td>1 (0.26%)</td>
</tr>
<tr>
<td>Nevi</td>
<td>0</td>
<td>0</td>
<td>34 (40.96%)</td>
<td>34 (8.83%)</td>
</tr>
<tr>
<td>p</td>
<td>0.000**</td>
<td>0.000**</td>
<td>0.651</td>
<td></td>
</tr>
</tbody>
</table>

From analyzing the results, we noticed that NMSC and MSC mostly occurred de novo (78.44%) (figure no. 2).

Figure no. 2. The correlation between preexisting lesions and cutaneous tumours

![Diagram showing correlation between pre-existing lesions and tumours.](image)

By conducting a comparative analysis of NMSC / MSC, we noticed that NMSC occurred more frequently on normal skin without preexisting lesions (82.59%), compared with MSCT (59.04%) (p = 0.006 **) (figure no. 3).

BCC occurred most frequently de novo (84.49%), followed by the appearance on the scars (8.16%) and actinic keratosis (7.35%), the results being statistically significant (p = 0.000 **).

SCC developed most frequently de novo (80.70%), followed by the appearance on cheilitis (8.77%) on actinic keratosis (7.03%), on scars and ulcers (1.75% ), the results being statistically significant (p = 0.000 **).

Our results reveal a roughly equal proportion of actinic keratosis converted into BCC (7.35%) and SCC (7.03%), in contrast to data found in the literature, where the malignant transformation of actinic keratoses in the SCC is higher than in BCC.(1)

MM appeared de novo in 59.04% of the cases, with the rest (40.96%) attributed to malignant transformations of the nevi lesions (p = 0.651) (table no. 1), the results being similar to those in medical literature.(2)

### DISCUSSIONS

Skin tumours can either arise de novo, on healthy-looking skin, or as a result of a malignant transformation of precancerous lesions or dysplastic nevi. In our study, skin cancers occurred predominantly de novo (78.44%), with the following distribution: 84.49% in NMSC and 59.04% in MSC.

In the literature, it was argued that the types of precancerous lesions which can set off NMSC are very diverse, including actinic keratoses and seborrheic, sebaceous nevus, basal cell nevi syndrome, epidermodysplasia verruciformis, vaccination scars, tattoos, chronic radiodermatitis, leukoplakia, erosive oral lichen planus, Bowen disease, eritroplazia, discoid lupus erythematosus (3), varicose ulcers, lupus vulgaris, necrobiosis lipoidica, sclerose-atrophic lichen, deep chronic fungal infections, epidermolysis bullosa (4,5), burns (6), osteomyelitis, acne conglobata, cellulite dissecans of the scalp, hidradenitis suppurativa, lymphogranuloma venereum etc.(7,8,9)

In this study, the most common pre-existing lesions in patients with BCC were scars (8.16%) and actinic keratoses (7.35%).

In patients with SCC, the most common precancerous lesions were cheilitis (8.77%) and actinic keratoses (7.05%). A special case worth noting was that of a malignant transformation into SCC of a mixed arterio-venous ulcer localized on the ankle that had been in the florid phase for several years. After three years since the amputation of the affected lower limb, the patient is alive without metastases. (Note: In our patient, this particular case was...)

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ulcer had first appeared 25 years ago, scarring repeatedly and presenting vicious epithelization). The studies published on the subject recognize that the transformation of chronic scarring in SCC may occur on average after 35 years (10), with a 30% rate of metastasis (10,11), and a 33% mortality rate.(12)

In a group of 200 patients with invasive SCC, the risk of metastasis was 12.5%, which was correlated with tumour size, grade of differentiation, perineural invasion and recurrence.(13)

MM may arise “de novo” in about 60% of cases, the rest arising from dysplastic nevi or nevi with a modified clinical character.(2) Patients with dysplastic nevi have a higher risk of developing MM during their lifetime, either through the evolution of a mole or directly on healthy skin (14); the presence of atypical nevi may be considered to be a marker for the identification of a category of patients with the relative risk for the development of MM.

In the group of patients with MM, 59.04% of the cases occurred de novo. This result is almost identical to the one from the literature.(2)

By analyzing the pre-existing lesions that turned malignant, we noticed that MSC developed upon clinically changed nevi (40.96%) and NMSC developed upon actinic keratoses, scars, cheilitis and mixed arteriovenous ulcers.

These results converge to the same idea: recognition of precancerous lesions and of dysplastic nevi, as well as establishing effective and early therapeutic measures could allow a significant decrease in the incidence of skin malignancies in the future and their diagnosis in the early stages of the disease. In this case, public education plays a very important role; the prospective patient should consult a dermatologist as soon as she identifies new skin lesions or any changes in nevi preexisting lesions. Unfortunately, in Romania patients delay consulting their doctor when faced with a suspicious lesion due to a scientifically unfounded fear of metastasis resulting from the tumour excision. Removing this fear will allow us to finally be aligned with developed countries, where patients seek endorsed medical advice in a much more timely manner.

In Australia, where the incidence of skin cancers, particularly MM, is high, the diagnosis is carried out to a large extent by the patient and the primary medical network, and it is then certified by dermatologists. This is reflected in the diagnosis of the disease in the early stages, a decrease of mortality in these tumours and a reduction of their treatment recurrence.(13)

CONCLUSIONS
1. In our group of patients, the clinical distribution on forms of tumour was: 63.64% BCC 21.56% MM and 14.80% SCC.
2. In our study, skin cancers occurred predominantly de novo (78.44%). Over two thirds of NMSC appeared on seemingly healthy skin (84.49%) and 59.04% of MSC arose de novo.
3. In the NMSC group that appeared on preexisting lesions, the most common pre-cancers were actinic keratoses (44.90%) and scarring (42.86%), followed by cheilitis (10.20%) and ulcers (2.04%).
4. The identification of pre-existing lesions, as well as a timely treatment, would reduce the incidence of both skin malignancies and mortality, especially in MSC.

REFERENCES