Contribution of Imaging Techniques in the Management of Cutaneous Pathology

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The management of skin pathology of tumors and inflammatory disorders is difficult due to the existence of constantly changing diagnostic and treatment algorithms. The incidence of cutaneous melanoma and other melanocytic or non-melanocytic tumors is currently increasing. Melanoma is difficult to treat in advanced stages due to its aggressiveness, and early diagnosis is required to improve the prognosis of these patients. Imaging techniques, such as classical and digital dermatoscopy can provide information on structure, vascular pattern, prognostic factors and detailed morphological analysis that can lead to improved individual management. This article presents a retrospective study that aims to analyze the contribution of imaging techniques to clinical and histological data.

Keywords: skin tumors, dermatoscopy, melanoma, melanocytic nevi

The management of tumor and inflammatory skin pathology falls within the area of interest in current scientific and applied scientific research due to the increased frequency of these diseases and the existence of constantly changing diagnostic and treatment algorithms. The incidence of skin tumors is steadily rising, with more than one patient diagnosed with other cancers in the last three decades [1]. In recent years, malignant melanoma and non-melanoma skin neoplasms have become an increasingly common form of cancer [2]. Melanoma accounts for less than 1% of the percentage of cutaneous tumors but accounts for the highest mortality rate, and therefore screening methods and diagnosis with high sensitivity and specificity are required to reduce mortality. Although the clear method of diagnosing tumor lesions is histopathological examination, non-invasive methods, such as digital dermatoscopy along with high frequency ultrasound, offer the possibility of increasing reliability of diagnosis correctness, establishing prognostic factors and therapeutic management with the possibility of avoiding invasive maneuvers [3-5]. Having been considered stethoscope of the dermatologist, dermatoscopy is used primarily to assess benign tumor lesions (melanocytic nevi, lentigo, seborrheic keratoses, dermatofibromas) and detect skin malignant lesions such as malignant melanoma, basal cell carcinomas or squamous cell carcinomas.

This article presents a retrospective study that aims to evaluate whether imaging investigations provide additional information that helps to increase diagnostic accuracy and the possibility of individualizing the treatment of skin lesions of the tumor type.

Experimental part

Materials and methods

The retrospective observational study was conducted after obtaining approval from the Institutional Ethics Commission. Patients between 18 to 75 years old with high-risk, intermediate or non-malignant pigmentary and non-pigmentary tumoral lesions were admitted at the St. Spiridon Hospital Dermatology Clinic and the offices in which doctors were working between 2015 and 2017. Excluded from the study were patients presenting neuropsychiatric pathology, deep neoplastic pathology, tumor lesions that had already undergone one or more surgical procedures, such as biopsy or excision, infected or with significant bleeding. A clinically integrative, dermatoscopic, histopathological analysis was performed and these data were introduced into the database and statistically analyzed using the Statistics 7 software.

Results and discussions

The study included at this stage a total of 38 patients diagnosed with pigmentary or non-pigmentary tumoral lesions. The following diagnostic imaging tests were used:

- Manual dermatoscopy: for the examination of tumor lesions, a non-polarized dermatoscope and / or dermatoscope with polarized light, with or without immersion fluid, were used depending on the type of tumor being analyzed.

- Videodermatostopy (digital dermatoscopy): A videodermatoscope was used to store images and assisted computer analysis by obtaining a DANAOS score for each tumor lesion.

- The histopathological examination is the gold standard diagnostic method in the evaluation of pigmentary and
non-pigmentary tumoral lesions, which analyzes in optical microscopy. Immunohistochemistry data are useful in the assessment of various pathologies as well as in cases of skin lesions where classical investigations do not provide conclusive data [6,7].

The analysis of the data shows that 63.1% of the registered patients were female and 36.9% male, with an average age of 41.05 years, ranging from 18 to 70 years. 89.46% of the patients came from an urban environment, and 34.3% of the patients said they had used photoprotection products during exposure to the sun. Patients reported a personal history of cardiovascular disease (hypertension, ischemic cardiopathy), vulgar psoriasis, autoimmune thyroiditis and operated breast cancer.

Figure 2 shows that the excised lesions had the following topography: 16% anterior thorax, 34% posterior thorax, 26% head and cervical region, 16% upper limbs, 8% lower limbs, with no mucosal involvement.

Table 1

| Diagnosis       | Tumors |                              |                  |                              |                  |
|-----------------|--------|------------------------------|------------------|------------------------------|------------------|
|                 |        | Benign                       | Suspected of malignancy | Malignant                    | Total            |
|                 |        | N                            | %                | N                            | %                |
| Clinical        |        | 29                           | 76.32            | 4                            | 10.53            | 5                | 13.16            | 38               |
| Histological    |        | 31                           | 81.58            | 1                            | 2.63             | 6                | 15.79            | 38               |

Comparison of clinical and histological diagnosis:

Table 2

| Clinical Diagnosis | Histologic Diagnosis | Total |
|--------------------|----------------------|-------|
| Positive           | Positive             | 10    |
|                    | Negative             | 28    |

Pigmented or non-pigmented malignant tumors, suspected tumors, as well as benign tumors which patients have requested excision for repeated trauma or aesthetic reasons, have been excised with surgical safety limits, and the samples were subsequently examined histopathologically using the Hematoxylin Eosin staining. If we consider that the histological examination provides the gold standard diagnosis, then we can compare the degree of correlation between the results of the clinical examination and the dermatoscopic examination with the results of the histological examination. Clinical and histological examinations revealed cases of benign tumors, cases of suspected malignant tumors and cases of malignant tumors, as presented in the table 1 and figure 3.

As can be seen, 10.53% of cases were diagnosed as suspected malignant by clinical examination and 2.63% of cases by histological examination, the difference being almost statistically significant at p <0.10: p = 0.05 > t = 1.389 > p = 0.10. Considering positive cases those cases diagnosed as malignant and those diagnosed as suspected of malignancy, we compared histological diagnosis with clinical and dermatoscopic diagnosis.
Comparison of dermatoscopic diagnosis assessed by malignant, suspected or benign tumor with histologic diagnosis

Dermatopyscopy is a non-invasive, practical and time-saving method that allows the diagnosis and evaluation of tumor morphological parameters that may or may not be detected in the clinical examination of cutaneous lesions, and studies show that this method improves the accuracy of clinical diagnosis by 20-30% [8].

In order to establish a dermatoscopic diagnosis, several diagnostic algorithms have been described over time. Most have the primary purpose of differentiating melanocytic lesions from non-melanocytic lesions. Subsequently, melanocytic lesions determine the benign or malignant nature. The ABCDE rule is one of the most common methods used in the clinical evaluation of pigment injuries. In this semiquantitative model, each of the four criteria is assigned a score, based on a calculation formula, the total dermatoscopic score (SDT) is determined. Other algorithms can be used such as:

a. The 7 improved criteria consisting of: atypical network, atypical vessels, white-blue veil (all valued at 2 points), regression structures, irregular dots / globules, irregular blotches - the presence of a single criterion is sufficient to perform a biopsy; and the presence of more than 2 points is equivalent to melanoma.

Digital dermatoscopy is a modern method of diagnosis and assessment of skin tumors, especially pigmented lesions, using video digital imaging, a powerful tool with the ability to memorize, analyze images and compare them over a period of time to track the evolution of cutaneous lesions. The images obtained from the videodermatoscopic examination can be analyzed by the clinician and can also be evaluated by assisted computer analysis system by calculating the DANAOS score based on the Artificial Intelligence principle. Digital dermatoscopy may allow early detection of malignant tumor lesions, the DANAOS score obtained from dermatoscopic analysis of tumors may range from 0 to 10, with values less than 4.75 being attributed to benign lesions, values greater than 5.45 for

| Statistic              | Value   | 95% Confidence Interval |
|------------------------|---------|-------------------------|
| Sensitivity            | 100%    | 63.05% to 100%          |
| Specificity            | 93.33%  | 77.93 to 99.18          |
| Positive Likelihood Ratio | 1.50   | 3.52 to 37.22           |
| Negative Likelihood Ratio | 0     |                         |
| Prevalence             | 21.05%  | 9.55% to 37.33%         |
| Positive Predictive value | 77.78% | 47.21% to 93.20%        |
| Negative Predictive Value | 100%   |                         |
| Accuracy               | 94.74%  | 82.25% to 99.36%        |

Table 3
RESULTS OF COMPARISON OF HISTOLOGICAL CLINICAL DIAGNOSIS

| Dermatoscopic Examination | Histologic examination | Total |
|---------------------------|------------------------|-------|
|                           | Positives | Negatives |       |
| Positives                 | 7 real positives | 2 false positives | 9 |
| Negatives                 | 1 false negative | 28 real negatives | 29 |
| Total                     | 8          | 30         | 38    |

Table 4
FREQUENCY OF CASES BY DERMATOSCOPIC DIAGNOSIS APPEARED THROUGH PRESENT / ABSENT MALIGNANT RISK AND HISTOLOGY

b. Pattern analysis - descriptive method.
c. The two-step algorithm: determining whether or not the tumor is melanocytic, and in the second stage establishing the type of tumor (nevus or melanoma).
d. The Menzies method consists of the presence of negative characteristics (symmetric pattern, the presence of a single color) and positive features of melanoma - in which the diagnosis of melanoma consists in the absence of negative characteristics and the presence of at least one positive characteristic.

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| Statistic              | Value   | 95% Confidence Interval |
|------------------------|---------|-------------------------|
| Sensitivity            | 87.3%   | 47.33% to 99.68%        |
| Specificity            | 93.33%  | 77.93 to 99.18          |
| Positive Likelihood Ratio | 13.13  | 3.55 to 51.36           |
| Negative Likelihood Ratio | 0.13   | 0.02 to 0.84            |
| Prevalence             | 21.05%  | 9.55% to 37.33%         |
| Positive Predictive value | 80%    | 11.18% to 93.83%        |
| Negative Predictive Value | 95.55% | 81.70% to 99.43%        |
| Accuracy               | 92.11%  | 78.62% to 98.34%        |

Table 5
RESULTS OF THE COMPARISON BETWEEN DERMATOSCOPIC DIAGNOSTIC APPRECIABLE BY MALIGNANT RISK / SUSPECTED LESION
The sensitivity of the digital dermatoscopy analysis was reported to be between 80% and 100% [10] and the specificity between 46 and 98% [11]. Similarly, in our study the sensitivity of the dermatoscopic diagnosis to the assessment of tumor malignancy / suspected tumor risk was 87.5%, \( t = 2.248 > p = 0.01 \) and the specificity was recorded as 93.33% (fig. 4).

Also, computer-assisted digital evaluation of skin tumor lesions provides the possibility of follow-up, particularly in patients with multiple atypical nevi [12], the dermatologist having the ability to compare the parameters of tumor lesions (maximum diameters, surface, asymmetry, margins, variety of color) at predetermined time intervals, as well as to analyze these lesions with the DANAOS score (dynamic values), which allows the possibility of establishing prognostic factors and the therapeutic plan (fig. 5).

The prognosis score of the DANAOS score was correlated with the low malignant dermatoscopic score with values lower than 4.75 in 28.94% of cases, values between 4.75 and 5.45 in 7.89% of cases and an increased risk of malignancy was attributed with values greater than 5.45 in 63.17% of cases.

The most common type of tumor analyzed was the compound melanocytic nevus (26.32%). The comparison of the frequency of the types of tumors diagnosed clinically and histopathologically revealed as shown in table 6 the following differences:

| Tumor type                        | Clinical                     | Histological                  | Diagnosed cases |
|-----------------------------------|------------------------------|--------------------------------|-----------------|
| Compound melanocytic nevus        | 19 26.32                     | 1 26.32                       | -               |
| Seborrheic verruca                | 6 15.79                      | 4 16.33                       | 2: dermal nevi  |
| Hemangioma                        | 2 5.26                       | 2 5.26                        |                 |
| Junctional melanocytic nevus      | 2 5.26                       | 2 5.26                        |                 |
| Dermal nevus                      | 3 7.89                       | 3 7.89                        |                 |
| Verruca vulgaris                  | 1 2.63                       | 1 2.63                        |                 |
| Dermatofibroma                    | 1 2.63                       | 1 2.63                        |                 |
| Blue Nevus                        | 3 7.89                       | 3 7.89                        |                 |
| Papilloma                         | 1 2.63                       | 1 2.63                        |                 |
| Dysplastic melanocytic nevus      | 4 16.33                      | 1 2.63                        | 2: nodular basal cell carcinoma; 1: blue nevus |
| Squamous cell carcinoma in situ   | 1 2.63                       | 1 2.63                        |                 |
| Nodular basal cell carcinoma      | 3 7.89                       | 1 2.63                        | 1: dermal nevus; 1: malignant melanoma |
| Malignant melanoma                | 1 2.63                       | 1 2.63                        |                 |

- from 6 cases diagnosed clinically with seborrheic wart, histopathologically there were 2 cases with non dermal nevi;
- from 4 clinically diagnosed cases with dysplastic melanocytic nevi, histopathologically 2 cases with nodular basal cell carcinoma and 1 case with blue nevi were diagnosed, the difference being significant at \( p <0.10; p = 0.05 > t = 1.389 > p = 0.10 \)
- from 3 clinically diagnosed cases with nodular basal cell carcinoma, histologically one case was diagnosed with non-dermal and 1 case with malignant melanoma.

Differences have not been statistically significant. Clinically, 5.27% of cases with benign tumors (76.32% compared to 71.05% histologically diagnosed) were diagnosed with 7.9% more cases of suspected malignant tumors (10.53% vs. 2. 63% histologically diagnosed, the
difference being almost statistically significant $p = 0.05$, $t = 1.389$ $p = 0.10$) and 5.27% more cases with malignant tumors (13.16% histologically diagnosed) (fig. 6).

The limits of this analysis may be given by the reduced number of patients, and data on the study of the effects of dermatoscopy on the current practice of the dermatologist are limited, requiring further research and prospective or retrospective studies in a larger number of patients.

Clarification:
The confidence intervals for sensitivity, specificity and accuracy are exact Clopper-Pearson confidence intervals.
The confidence intervals for Likelihood Ratios are calculated using the Log method.

NOTE
If the prevalence of the disease is known, then the positive and negative predictive value can be calculated using a formula based on Bayes’ theorem:

\[
PPV = \frac{\text{sensitivity} \times \text{prevalence}}{\text{sensitivity} \times \text{prevalence} + (1 - \text{specificity}) \times (1 - \text{prevalence})}
\]

And respectively:

\[
NPV = \frac{\text{specificity} \times (1 - \text{prevalence})}{(1 - \text{specificity}) \times \text{prevalence} + \text{specificity} \times (1 - \text{prevalence})}
\]

Conclusions
Taking into account the results obtained in the study, it is observed that the specificity of the dermatoscopic examination with the assessment of the malignant or suspected risk of tumor lesions is comparable to that of the clinical examination with the naked eye. It is also dependent on the examiner and needs a dermatoscopic analysis training to increase sensitivity and diagnostic specificity. It is also noted that videodermatoscopic evaluation with lesion analysis and DANAOS score calculation can not replace clinical examination, but in the case of atypical tumor lesions, this method can give the dermatologist a quick second expert opinion as literature data [13,14]. Melanoma can be clinically and sometimes even dermatoscopically not differentiated from benign lesions, representing a challenge for the dermatologist in the formulation of the diagnosis until the histopathological confirmation.

References
1. STERN RS. Prevalence of a history of skin cancer in 2007: results of an incidence-based model. Arch Dermatol 2010; 146(3):279-282.
2. GHEUCĂ SOLOVASTRU L, VAĂ D, STÎNCANU A, CIUBĂRA AM, ANDREȘE E. The psychosocial impact on patients with skin neoplasia.Bulletin of Integrative Psychiatry2013; 3(58): 33-38.
3. SCHMIDT-WENDTNER MH, Br J Dermatol Dill-Muller D. Ultrasound technology in dermatology. SeminCutanMedSurg 2008; 27: 44-51.
4. GUITERA P, LILX, CROTTY K ET AL. Melanoma histological Breslow thickness predicted by 75-MHz ultrasonography. Br J Dermatol 2008; 159:364-369.
5. CRISAN, D., BADEA, A.F., CRISAN, M., RASTIAN I., SOLOVASTRU GHEUCĂ, L., Integrative analysis of cutaneous skin tumours using ultrasonographic criteria. Preliminary results. Med Ultrason 2014;16(4):285-290.
6. SALAHORU, P., GHICIUC, C.M., GRIGORESCU, C., HINGANU, M.V., LUPUSORU, C.E., Use of immunochemistry to establish etiology in the case of primary spontaneous pneumothorax patients. Rev. Chim. (Bucharest), 69, no. 7, 2018, p. 2251-2253.
7. GRIGORESCU, C., GAVRIL, L.C., GAVRIL, L., LINGULEAC, T., CIUNTU, B.M., PATRASCU, A., SALAHORU, P., Anti-receptor Concentration And Over The Clinical Manifestations On Myasthenia Gravis Patients. Rev. Chim (Bucharest), 69, no. 9, 2018, p. 2591-2593.
8. MASOOMEH B, HAIEDEH G, PARISA M, ARASH T, ZAHRĂ S, MASOOD A. Computer-aided dermoscopy for diagnosis of melanoma. BMC Dermatol 2005; 5:8.
9. SERRAOV, BAPTISTA J, PARIS F, FERREIRA A. Digital dermoscopy. Review of 652 lesions analysed by the DANAOS system. Skin Cancer 2006; 21(4):185-198.
10. ERCAL F, CHAWLA A, STOECKER WV, LEE HC, MOSS RH. Neural network diagnosis of malignant melanoma from color images. IEEE Trans Biomed Eng1994; 41(9):837-45.
11. FARINA B, BARTOLI C, BONO A, COLOMBO A, LUALDI M, TRAGNI G, MARCHESINI R. Multispectral imaging approach in the diagnosis of cutaneous melanoma: potentiality and limits. Phys Med Biol 2000; 45(5):1243-54.
12. GREEN A, MARTIN N, PFITZNER J ET AL. Computer image analysis of melanoma in the diagnosis of melanoma. J Am AcadDermatol1994; 31: 958–64.
13. SCHWARZER G, VACH W, SCHUHMACHER M. On the misuses of artificial neural networks for prognostic and diagnostic classification in oncology. Stat Med 2000; 19: 541-61.
14. LISBOA PJ. A review of evidence of health benefit from artificial neural networks in medical intervention.Neural Netw. 2002;15(1):11-39.

Manuscris received: 6.08.2018