Integrated care pathways and the hub-and-spoke model for the management of non-melanoma skin cancer: A proposal of the Italian Association of Hospital Dermatologists (ADOI)

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Abstract

The term non-melanoma skin cancer (NMSC) refers to skin cancer different from melanoma, and it is usually restricted to basal cell carcinoma (BCC), squamous cell carcinoma (SCC) and their pre-cancerous lesions, e.g., actinic keratosis. These conditions represent the most frequent tumors in Caucasians and are characterized by an increasing incidence worldwide and a high socio-economic impact. The term Integrated Care Pathway (ICP) refers to “a complex intervention for the mutual decision making and organization of care processes for a well-defined group of patients during a well-defined period”. The purpose of this paper is to present a proposal from the Italian Association of Hospital Dermatologists (ADOI) for an ICP organization of care of NMSC, considering the hub-and-spoke model in the different geographical areas.

This proposal is based on the most recent literature and on documents from the Italian Association of Medical Oncology (AIOM), the European consensus-based interdisciplinary guidelines from the European Association of Dermato-Oncology (EADO), and the National Comprehensive Cancer Network (NCCN).

We initially discuss the NMSC outpatient clinic, the role of the multidisciplinary working groups, and the hub-and-spoke model regarding this topic. Then, we define the ICP processes specific for BCC and SCC.

The ICP for NMSC is an innovative strategy to guarantee the highest possible quality of health care while the hub-and-spoke model is crucial for the organization of different health care structures. Considering the importance on this topic, it is essential to create a valid ICP together with an efficient organization within the different geographical areas.

Introduction

The term non-melanoma skin cancer (NMSC) traditionally comprises skin cancer that arise from keratinocytes of the epidermis and mainly includes basal cell carcinoma (BCC), squamous cell carcinoma (SCC) and actinic keratosis (AK). The incidence of NMSC is constantly growing worldwide and, therefore, the management and treatment of the aforementioned neoplasms has become an important challenge for the health system.1

The cellular origin of BCC has not been completely elucidated but it is thought to arise from follicular and interfollicular epidermal basal keratinocytes.2 BCC is predominantly locally invasive and rarely metastasizes (0.05-0.1% of cases).3 The incidence of a first BCC in Italy has been estimated at about 87.6 per 100,000 inhabitants/year and represents 15% of all cancers.4

SCC originates from the squamous cells of the epidermis. SCC includes “invasive” and other “non-invasive” forms such as keratoacanthoma, Bowen’s disease, Bowenoid papulosis and Queyrat’s erythroplasia.3 SCC can arise from a previous AK or de novo and is more often localized in photo-exposed areas but can also affect the mucous membranes and genitalia. SCC represents the second most frequent skin cancer (20%), after BCC, with an incidence of about 28.9 per 100,000 inhabitants/year. In 1-5% of cases, it can give distant metastases and such metastases are associated with an average survival of two years.3

The main risk factors related to the onset of NMSC are exposure to UV radiation, advanced age, light skin type, chronic immunosuppression, ionizing radiation, arsenic exposure, human papillomavirus infections, burns, chronic inflammatory processes, and specific genodermatoses.3,5

The Integrated Care Pathway (ICP) is, according to the definition adopted by the European Pathway Association (EPA), “a complex intervention for the mutual decision making and organization of care processes for a well-defined group of patients during a well-defined period”.6 The aim of an ICP is to enhance the quality of care across the continuum by improving risk-adjusted patient outcomes, promoting patient safety, increasing patient satisfaction, and optimizing the use of resources.6

The ICP of the patient affected by NMSC is multidisciplinary. A multidi-
The increasing number of NMSC, the severity of some cases and the possibility to utilize recent treatments for advanced or metastatic cases of BCC (i.e., sonidegib and vismodegib) and SCC (i.e., cemiplimab) require an integration of community-based health care services with specialized hospital clinics that provide the optimal environment to address the complex needs of the cases and improve outcomes. This approach requires coordination across different levels and sites of care within and beyond the health sector. The hub-and-spoke organization design is a model characterized by service delivery assets into a network involving an anchor establishment (hub) offering a full array of services, complemented by secondary establishments (spokes) offering more limited service arrays, routing patients needing more intensive services to the hub for treatment (Figure 1). The hub-and-spoke model favours a healthcare network involving a main campus and one or more satellite campuses and this model is more efficient than organization designs organized in multiple sites. Hub-and-spoke network structure could vary with satellites that could be increased as needed or desired. If geographic distance makes satellite-to-hub access difficult, an extra hub could be added. Therefore, the hub-and-spoke model represents an organization of care that works collaboratively with the primary care sector and is greatly integrated with community-based multidisciplinary teams of health care professionals and specialty care. Strategic centralization brings many benefits to the hub-and-spoke organization design. Telemedicine, particularly teledermatology (TD), represent a valid and well-established tool to help to coordinate the hub-and-spoke workflow. TD has been proven in various studies to be a valid triage system for skin cancer detection.

**Integrated care pathway for non-melanoma skin cancer**

The ICP for NMSC should consider different institutional levels and the risk of the BCC or SCC as shown in Figure 2 (Tables 1 and 2 for BCC and SCC risk classification). Local dermatology outpatient clinic should treat patients affected by low risk BCC or SCC, while, general hospital with a Dermatology Unit could manage intermediate and high-risk BCC or high and very high-risk SCC. More complicated cases (i.e., locally advanced or metastatic BCC or SCC, high risk BCC, or very high-risk SCC) should be managed by tertiary node with skin cancer centres and oncologic services organised in multidisciplinary working group (MWG). The NMSC outpatient clinic is the clinic in which the patient suffering from this neoplasm is checked by a specialist dermatologist in order to treat and follow-up the patient. This dermatological outpatient clinic is configured as a central part in the ICP process for NMSC because it is dedicated to the diagnosis, management and follow-up of patients affected by most of these tumors. The MWG is essential for locally advanced forms of BCC and/or SCC for which there is no indication for surgery or radiotherapy or for metastatic NMSC (Figure 3). The MWG has been introduced to reduce the variation in decision-making between different specialists and for patients and their carers to access the best care possible. Considering the ageing population with multiple comorbidities, the number of treatment options becoming available, and the complexity of some NMSC, the MWG is crucial in order to offer the gold standard oncologic services. The MWG should periodically meet to discuss specific NMSC cases. In such meetings, clinicians should select appropriate cases to be discussed and these cases should be treated following specific ICP for NMSC, if possible. Routine cases should not be discussed in MWG meetings and should be treated as per protocol. Finally, the MWG meetings offer a source of support, education and management updates for the clinicians and trainees in a constantly and rapidly changing area. Clinical diagnosis of NMSC could be enough in many cases but the use of dermoscopy may improve diag-
nostic accuracy. In the diagnostic process, it is important to consider medical history (advanced age, higher phototype, previous diagnosis of NMSC, excessive sun exposure, immunosuppression therapy) and physical examination (location, size, infiltration and margins of the lesion). In case of diagnostic doubt, it is essential to perform a skin biopsy and histological examination.

Surgery represents the first-line therapy for BCC and SCC because it allows histopathological analysis and low risk of relapsing. Other alternative techniques are indicated in case of low-risk BCC or in case of in situ SCC and are discussed below.

The histological examination reports the local staging (pT) following the excisional biopsy of the NMSC which allows to define and plan any subsequent instrumental or surgical procedures. Furthermore, it must include the following information: patient data, report number/year, report date, site of tumor, macroscopic finding, histological subtype (in case of high-risk neoplasm), histological grade (for SCC: well differentiated - G1 moderately differentiated - G2, poorly differentiated - G3), state of lateral and deep margins / complete excision, tumor thickness (related to NMSC), possible perineural invasion and lymphatic / vascular invasion.

Any instrumental examinations will be planned based on the type of NMSC and the clinical assessment. These examinations include ultrasound of the loco-regional lymph nodes which is indicated in case of high-risk invasive SCC. Advanced forms of SCC or BCC may require the use of diagnostic imaging techniques such as computed tomography and magnetic resonance imaging that allow evaluation of the local extension of the neoplasm, the infiltration of adjacent anatomical areas and the possible presence of nodal or distant organ metastasis.

**Basal cell carcinoma**

**Risk stratification for basal cell carcinoma**

BCC can be classified into low-risk and high-risk according to prognostic factors such as tumor size, definition of clinical margins (poorly-defined lesions are at higher risk), histological subtype (morphoeform, and metatypical BCC represent high-risk lesions), histological features (perineural and/or perivascular invasion is a marker of higher risk), recurrence, and tumor location (Table 1). With regards to the location, high-risk zones are the nose, periorificial areas of the head and neck; intermediate-risk zones are the forehead, cheek, chin, scalp, and neck; low-risk zones are the trunk and limbs. Low-risk BCCs are superficial BCC, Pinkus tumor, and small nodular BCC on intermediate or low-risk areas while high-risk BCCs present at least one poor prognostic factor (Table 1). Furthermore, French guidelines also added an intermediate-risk to classify recurrent superficial BCC from other recurrent BCC, and some nodular BCCs according to size and location (Table 1). This classification has been used by the most recent AIOM guidelines regarding BCC (2020).

**Therapy of basal cell carcinoma**

The first-line therapy for BCC is surgery because it allows the complete excision of the skin cancer and the preservation of the cosmetic and functional aspects. An evidence-based review regarding the interventions for BCC reported that the best results have been obtained with surgery.
Other approaches indicated below could be used mainly in case of low-risk BCC.

**Surgery and Mohs micrographic surgery**

Standard excision is the primary treatment of BCC while the Mohs technique, not always feasible in medical institutions, could be preferable for the high-risk BCC. Standard excision with postoperative margin assessment (SEPM) must guarantee a post-operative histological evaluation with free margins. Regarding the extension of free margins, the National Comprehensive Cancer Network (NCCN) recommends clinical margins of at least 4 mm for low-risk BCC treated with SEPM. Indeed, Brodland et al. showed that for well-circumscribed BCC <2 cm in diameter, excision with 4-mm clinical margins guarantees a complete removal in more than 95% of cases. Furthermore, in case of SEPM for high-risk BCC, wider surgical margins compared to low-risk BCC are needed. A greater recurrence rate is predictable in case of high-risk BCC (Table 1).

MMS (also known as chemosurgery, microscopically controlled excision, or histographic surgery) is a surgical approach consisting in a complete excision of the tumor followed by an examination of the microscopic margins. This technique could be considered as the treatment of choice for high-risk and recurrent BCCs because it is always feasible in medical institutions, could be preferable for the high-risk BCC.

<table>
<thead>
<tr>
<th>Low-Risk</th>
<th>Intermediate-risk</th>
<th>High-Risk</th>
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</thead>
<tbody>
<tr>
<td>Prognostic groups for BCC</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Superficial primary BCC</td>
<td>Superficial recurrent BCC</td>
<td>Morpheaform or poor-defined types</td>
</tr>
<tr>
<td>Nodular primary BCC when:</td>
<td>Nodular primary BCC when:</td>
<td>Nodular primary BCC when:</td>
</tr>
<tr>
<td>1 cm in intermediate risk area*</td>
<td>&gt;1 cm in intermediate risk area*</td>
<td>&gt;1 cm in high risk area*</td>
</tr>
<tr>
<td>&lt;2 cm in low risk area*</td>
<td>&gt;2 cm in low risk area*</td>
<td>Recurrent forms (apart from superficial BCC)</td>
</tr>
<tr>
<td>Pinkus tumor</td>
<td>&lt;1 cm in high risk area*</td>
<td>Histological forms: aggressive,*</td>
</tr>
<tr>
<td>Therapeutic strategies</td>
<td>Surgery and Mohs micrographic surgery</td>
<td>Sonidegib</td>
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<tr>
<td>Suspicous superficial BCC, Surgery</td>
<td>Mohs micrographic surgery</td>
<td>Sonidegib</td>
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<tr>
<td>Defined superficial BCC: surgery, topical therapies, local therapies, PDT</td>
<td>Mohs micrographic surgery</td>
<td>Sonidegib</td>
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<tr>
<td>Nodular BCC surgery</td>
<td>Mohs micrographic surgery</td>
<td>Sonidegib</td>
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</table>

*High-risk zones are the nose, periorbital areas of the head and neck; intermediate-risk zones are the forehead, cheek, chin, scalp, and neck; low-risk zones are the trunk and limbs. Aggressive histological forms include micronodular, morpheaform, and metatypical basosquamous forms. Perineural invasion also seems to be a histological sign of aggressiveness.

### Table 1. Prognostic groups for BCC and therapeutic strategies (Trakatelli et al. 2014; Dandurand 2006). BCC can be classified into low-risk, intermediate-risk, and high-risk according to prognostic factors as indicated by the AIOM guidelines on BCC (2020). Therapeutic strategies are indicated into three main groups of BCC: low-risk, intermediate and high-risk, and locally advanced.

<table>
<thead>
<tr>
<th>Risk group</th>
<th>Low-risk</th>
<th>Intermediate-risk</th>
<th>High-risk</th>
</tr>
</thead>
<tbody>
<tr>
<td>Location/size</td>
<td>Trunk, extremities &lt; 2 cm</td>
<td>Trunk, extremities &gt; 2 cm &lt; 4 cm</td>
<td>≥ 4 cm</td>
</tr>
<tr>
<td>Borders</td>
<td>Well defined</td>
<td>Poorly defined</td>
<td></td>
</tr>
<tr>
<td>Primary vs recurrent</td>
<td>Primary</td>
<td>Recurrent</td>
<td></td>
</tr>
<tr>
<td>Immunosuppression</td>
<td>(-)</td>
<td>(+)</td>
<td></td>
</tr>
<tr>
<td>Site of prior RT or chronic inflammatory process</td>
<td>(-)</td>
<td>(+)</td>
<td></td>
</tr>
<tr>
<td>Rapidly growing tumor</td>
<td>(-)</td>
<td>(+)</td>
<td></td>
</tr>
<tr>
<td>Neurologic symptoms</td>
<td>(-)</td>
<td>(+)</td>
<td></td>
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</tbody>
</table>

**Histopathology**

<table>
<thead>
<tr>
<th>Degree of differentiation</th>
<th>Well or moderated differentiation</th>
<th>Poorly differentiation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Adenoid, adenosquamous, or metaplastic subtypes</td>
<td>(-)</td>
<td>(+)</td>
</tr>
<tr>
<td>Depth thickness or level of invasion of subcutaneous fat</td>
<td>≤ 6 mm or no invasion beyond subcutaneous fat</td>
<td>&gt; 6 mm or invasion</td>
</tr>
<tr>
<td>Perineural involvement</td>
<td>(-)</td>
<td>(+)</td>
</tr>
<tr>
<td>Lymphatic or vascular involvement</td>
<td>(-)</td>
<td>(-)</td>
</tr>
</tbody>
</table>

**Low risk**

<table>
<thead>
<tr>
<th>Therapeutic strategies</th>
<th>Surgery</th>
</tr>
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<tbody>
<tr>
<td>Mohs micrographic surgery</td>
<td></td>
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**High-risk and very high-risk**

<table>
<thead>
<tr>
<th>Therapeutic strategies</th>
<th>Surgery</th>
</tr>
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<tbody>
<tr>
<td>Cemiplimab</td>
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**Locally advanced or metastatic**

<table>
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<tr>
<th>Therapeutic strategies</th>
<th>Surgery</th>
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</table>

*Risk stratification should be based on the highest risk factor present. Preoperative clinical tumor diameter. If clinical evaluation of incisional biopsy suggests that microstaging is inadequate, consider narrow margin excisional biopsy. Deep invasion is defined as invasion beyond the subcutaneous fat OR ≥ 6 mm. Location on the head, neck, hands, feet, pretilia, or anogenital area constitutes high risk based on location, independent of size.
showed a higher long-term cure rates compared to other surgical approaches and a maximum preservation of normal tissue in relation to conventional surgery. It has been reported that 5-year recurrence rates for primary and recurrent BCCs treated with MMS are 1% and 5.6%, respectively, compared with 10.1% and 17.4%, respectively, for SEPM. Unfortunately, limited medical institutions in Italy have all the necessary facilities to implement this specific practice.

Furthermore, it could be possible to consider the excision with complete circumferential peripheral and deep margin assessment (CCPDMA), that is an advanced surgical treatment using intraoperative frozen section assessment of all deep and peripheral margins, as an efficient alternative to MMS.12

Local therapy with curettage, electrodesiccation, cryosurgery and laser therapy

These therapies must be limited to low-risk BCCs because generally they lack histological examination. Furthermore, in case of doubt regarding the diagnosis, an incisional biopsy must be performed before the treatment is performed.

Curettage and electrodesiccation are considered as fast and cost-effective technique, as indicated by the NCCN, for selected low-risk BCCs.12

Cryosurgery is an easy, fast, and cost-effective technique able to destroy the skin cancer by freeze-thaw cycles. Some large case series report cure rates from 94% to 99% for BCC.23

Laser therapy has been used for BCC treatment as monotherapy and adjunct therapy.24 The most utilized lasers reported are superpulsed carbon dioxide and pulsed neodymium-based laser therapy.25,26 Reactive hyperemia, edema, scarring, and soreness could occur as adverse effects (AEs) of these therapies.3

Topical therapies

Imiquimod 5% cream and 5-fluorouracil (5-FU) 5% cream are used for the treatment of superficial BCC but their use should be limited to BCCs that could not be treated with other regimens or when cosmetic results are of major concern.27,28

Imiquimod 5% cream is an immune response modifier approved for the treatment of non-facial superficial BCC. Treatment regimen for BCC is five times weekly for six weeks. In a randomized controlled trial (RCT) that used twice daily imiquimod 5% for 12 weeks for superficial BCC, a 100% histologic clearance after 6 weeks of treatment has been reported.29 It has been shown that the application of imiquimod 5% cream once daily for 12 weeks for nodular BCCs led to a 76% clinical clearance.30 Imiquimod 5% cream is also used in case of basal cell nevus syndrome or Gorlin-Goltz syndrome.31 AEs of imiquimod 5% cream are local erythema and irritation of the skin and systemic effects include fatigue, fever and exfoliative dermatitis.

5-FU is recommended for superficial BCC but not for nodular forms.3 A statistically equivalent efficacy between 5-FU and imiquimod 5% in treating superficial BCC at a 12-month follow-up has been reported in a RCT.32 Other studies with longer follow-up showed a superiority of imiquimod compared to 5-FU.33 AEs of 5-FU could be erythema, swelling, and erosions.34

Photodynamic therapy

Photodynamic therapy (PDT) is a technique to mainly treat superficial BCCs or thinner nodular subtype, generally in patients affected by extensive or multifocal disease or multiple AKs.3 It consists in the application of a photosensitizing agent, aminolevulinic acid or methyl aminolevulinate, followed by irradiation with a light source. Studies reported that cure rates for BCC range from 70% to 90% but some of these studies had short follow-up periods.35 The PDT treatment protocol for BCC generally consists of two separate sessions, interrupted by a week, repeatable after three months in case of recurrence of the cancer.

Intralresional therapy

Intralresional chemotherapy is rarely used and mainly for high-risk BCC in patients not candidates for surgical therapy. It consists in the application of drugs such as 5-FU, interferons, interleukin-2, and bleomycin, and it has been used to treat BCC with variable results. AEs are unusual and mainly dose-dependent consisting of local effects at the treatment site and flu-like symptoms.36,37

Elettrochemotherapy

Elettrochemotherapy (ECT) is a cancer therapy that combines the administration of a chemotherapy agent to the delivery of permeabilizing pulses released singularly or as bursts. It is based on the principle of electrification of the cell membrane. This therapy is used for the local treatment of cutaneous metastases or for the treatment of primary cutaneous tumors such as BCC in patients not candidates for surgical therapy. In a single-center study, the complete response rate for 84 BCC patients, ineligible for conventional treatments, treated with ECT was overall 50%.38
target genes include GLI1, PTCH1 and HH interacting protein (HHIP1) that regulate the pathway itself. The outcome of the HH signaling depends on transcription of several cell-specific targets mediating different cellular responses, including proliferation, differentiation, cell survival, self-renewal, angiogenesis, epithelial-mesenchymal transition. Aberrant activation of HH pathway is a tumor-driver in BCC pathogenesis. Therefore, the inhibition of the HH pathway can be considered a valid strategy to counteract neoplastic growth. The most frequent alterations in BCCs are the loss or inactivating mutations in PTCH1 or SUFU, as well as activating mutations in SMO or GLI. HH pathway activity can be inhibited through several mechanisms, including inhibition of the receptor-ligand interaction, direct binding to SMO, and inhibition of GLI transcription factors. Currently, in Italy, two targeted drugs are indicated and approved for the treatment of advanced BCC: vismodegib and sonidegib. The mechanism of action of both drugs consists in the inhibition of SMO and, in turn, in the prevention of the signal cascade, maintaining the suppression of the transcription factors GLI. Thus, the two molecules have both a cytostatic and cytotoxic action on tumoral cells.

Vismodegib is indicated in both locally advanced and metastatic BCC. Historically, it was the first member of the HH pathway inhibitors (HPIs) class that is now considered to be a first-line treatment option. The drug is administrated orally (150mg/d), every day at the same time. No dose reductions are provided for in the technical data sheet.

The approval of vismodegib by Food and Drug Administration (FDA) and by the European Medicines Agency (EMA) in 2013 was based on the results of a phase II, open-label, noncomparative, international trial (ERVANCE BCC), which showed high rates of tumor control in the indicated patient populations, including individuals with or without Gorlin syndrome. The pivotal ERVANCE BCC study had enrolled 104 patients with locally advanced and metastatic BCC. The primary endpoint of the study was the ORR (Objective Response Rate), assessed through RECIST Criteria (Central Review), and results indicated 47.6% for locally advanced BCC and 33.3% for metastatic BCC. When assessed through RECIST criteria and Investigator Review, the ORR was 60.3% for locally advanced BCC and 48.5% metastatic BCC. The long-term update of the study demonstrated the durability of the response, the efficacy and the long-term safety. However, adverse events were significant. Indeed, discontinuation rate due to adverse effects was 21.2%. The deaths were 31.7% but none were related to vismodegib. The observational open-label STEVIE study was aimed at assessing the safety profile of vismodegib as primary endpoint in a more representative population. Thus, 1215 patients were enrolled. Most patients showed treatment-related side effects, including muscle spasms, alopecia, dysgeusia, weight loss, and asthma. Secondary endpoint was efficacy. The ORR assessed through RECIST 1.1 Criteria and Investigator Review was 68.5% for locally advanced BCC and 36.9% for metastatic BCC. Primary and secondary resistance to vismodegib has been reported, albeit at a low rate compared with some other targeted therapies. Vismodegib is therefore an effective and generally well tolerated systemic therapy and, since its regulatory approval, has become an established treatment option in clinical practice for patients with locally advanced and metastatic BCC that can no longer be suitably controlled with surgery and/or radiotherapy. However, some limitations of vismodegib treatment should be kept in mind. The inevitably occurring side effects of vismodegib lead to a significant rate of treatment discontinuation limiting overall drug exposure. Hence, long-term continuous treatment with vismodegib is not feasible in most patients.

Sonidegib is indicated in locally advanced BCC. The drug is administrated orally (200mg/d), every day at the same time, away from food.

The results of the pivotal multicenter phase II study (BOLT) which evaluated efficacy and safety of Sonidegib, led to the approval of the drug as a first-line treatment for locally advanced BCC. This trial enrolled 230 patients with advanced BCC and compared two dosing regimens (200 vs. 800 mg per day) of Sonidegib in a double-blinded, 1:2 randomized fashion. Sonidegib 200 mg demonstrated a better safety-risk profile than 800 mg at 30 months, with lower rates of grade 3/4 adverse events (43.0% vs. 64.0%) and adverse events leading to discontinuation (30.4% vs. 40.0%). Treatment-related side effects included muscle spasms, alopecia, dysgeusia, weight loss, and asthenia. Adverse events were managed with dose adjustments or interruptions, since Sonidegib offers in label the option for dose reduction of the drug. Patients receiving 200 mg of therapy had an ORR assessed through the stringent mRECIST criteria of 56.1% and of 71.2% assessed through central and investigator review respectively for locally advanced BCC, with a mDOR and a PFS of 26.1 and 22.1 months respectively (central review). On the contrary, objective response rates with 200mg were 7.7% (Central Review) and 23.1% (Investigator Review) in metastatic BCC.

A group of clinical experts in the management of locally advanced BCC summarized in a recent paper the clinical and pharmacological profiles of sonidegib and vismodegib based on published data and their own clinical experience. They highlighted that one key difference between the two pivotal studies was the criteria used to assess BCC severity. ERIVANCE used the conventional Response Evaluation Criteria in Solid Tumors (RECIST), while the more recent double-blind randomized BOLT trial used the more stringent modified RECIST (mRECIST). A preplanned analysis presented the outcomes from BOLT with RECIST-like criteria, and this enabled the experts to discuss relative efficacy outcomes for the two treatments. Centrally reviewed objective response rate (ORR) for vismodegib was 47.6% (95% CI: 35.5-60.6) at 21-month follow-up using RECIST. Using RECIST-like criteria, the ORR for sonidegib according to central review at 18-month follow-up was higher, at 60.6% (95% CI: 47.8-72.4). Both treatments were associated with similar patterns of adverse events. However, sonidegib demonstrated a longer time to adverse events onset (except for fatigue), with less frequent and less severe adverse events compared with vismodegib.

The pharmacokinetic profile of sonidegib and vismodegib shows several differences, such as volume of distribution and half-life. The consensus among the experts is that these pharmacokinetic differences could related to the differences seen in tolerability and efficacy between the two drugs. Although the efficacy and side effect profile of Sonidegib trial appear generally comparable to results from large-scale studies with vismodegib for locally advanced BCC, direct comparative clinical studies would be necessary to thoroughly assess differences. Therefore, currently the only drug approved for the treatment of metastatic BCC is vismodegib whereas, for locally advanced BCC, two aforementioned therapeutic possibilities are possible.

Next landscape in advanced systemic treatment in case of progression or intolerance to HH inhibitors

Cemiplimab-rwlc is an anti-PD-1 antibody approved for treatment of advanced cutaneous SCC. On February 9, 2021, the Food and Drug Administration (FDA) granted regular approval to cemiplimab-rwlc for patients with locally advanced BCC previously treated with a HH inhibitor or for whom a HH inhibitor is not appropriate. Furthermore, FDA granted accelerated approval of the drug as a first-line treatment for advanced SCC.
approval to cemiplimab-rwlc for patients with metastatic BCC previously treated with a HH inhibitors or for whom a HH inhibitors is not appropriate. Recently has been reported an open-label, multicentre, single-arm, phase 2 trial across 38 outpatients clinics, that analyzed the data of cemiplimab in patients with locally advanced BCC after HP inhibitor therapy. An OR per independent central review was observed in 26 (31%; 95% CI 21-42) of 84 patients, confirming that cemiplimab exhibited clinically meaningful anti-tumor activity in patients with locally advanced BCC after HH inhibitor therapy. Based on these data, on May 2021 the Committee for Medicinal Products for Human Use (CHMP) adopted a positive opinion regarding cemiplimab indicated, as monotherapy, for the treatment of adult patients with locally advanced or metastatic BCC who have progressed on or are intolerant to a HH inhibitor.

Squamous cell carcinoma

Risk stratification for squamous cell carcinoma

A risk assessment for SCC has to be made to determine the treatment and follow-up for patients. Table 2 shows SCC risk stratification according to the most recent NCCN Guidelines considering SCC risk factors associated with recurrence and metastasis. Risk factors for SCC stratification include location and size of the tumor, primary versus recurrent disease, immunosuppression of the patient, site of prior RT or chronic inflammatory process, neurologic symptoms, and histopathological features (Table 2). Concerning the actual histopathological staging systems, it is worth to notice that the American Joint Committee on Cancer (AJCC) 8th edition SCC staging system expanded the criteria for upstaging to T3 and also included extranodal extension as a risk factor for upstaging of the N classification (Tables 3 and 4). Otherwise, the AJCC 8th edition is limited to head and neck tumors. An alternative T staging system (the Brigham and Women’s Hospital [BWH] Tumor Staging for SCC) proved in its ability to stratify low-risk versus high-risk tumors and it can be applied to tumors across all body sites.

Therapy of squamous cell carcinoma

The primary goal for SCC therapy is the complete removal of the skin cancer with the maximal functional and cosmetic preservation. Surgical excision alone guarantees a successful treatment for SCC with a good prognosis and cure rates greater than 90%. Besides surgical therapy, traditional techniques such as curettage, electrodessication, cryosurgery, lasertheraphy, and PDT are available for non-invasive SCC forms like Bowen disease. Furthermore, RT is a valid and curative treatment strategy for SCC.

Surgery and Mohs micrographic surgery

Traditional surgery and MMS are the two different surgical approaches that may be utilized in patients with primary SCC. It has been reported that SEPMA guarantees a 5-year disease-free rates of 91% or higher for SCC. Safety excision margins have to be defined according to the risk of subclinical extensions, recurrence or and metastasis of the skin cancer depending on the low or high-risk factors for SCC. Considering low-risk SCC, AION and NCCN guidelines recommend free-margins of at least 4-mm for low-risk types treated with SEPMA. The European consensus group proposed a 5-mm margin for low-risk SCC. Otherwise, for high-risk SCC, NCCN guidelines recommend wider surgical margins compared to low-risk SCC and postoperative margin assessment. NCCN guidelines does not recommend a defined margin for standard excision for high-risk SCC due to the wide variability of clinical characteristics that defined this type of SCC. The European consensus group recommends 6-10 mm safety margins for high-risk SCC. In case of positive margins, it should be performed a re-excision, for operable cases. In case of margins that appear more limited than the recommended safety margins due to the tissue shrinkage a wider excision should be considered. We also recommend, as alternative to surgery, a close follow-up.

MMS offers the highest rate of R0 resection (i.e., no cancer cells seen microscopically at the primary tumor site), above 90%, with lower recurrence rates (0-4%) compared to traditional surgery (3.1-8.0%). MMS is mainly considered for patients with high-risk SCC to obtain a complete tumor resection with optimal anatomic and functional preservation. Otherwise, MMS is performed in limited centers and more time-consuming, labour-intensive, and expensive compared to traditional surgery. Van Lee et al., confirmed in a retrospective study that MMS could be superior to standard excision for SCC of the head and neck for the lower rate of recurrence.

Alternative treatment for low-risk squamous cell carcinoma: curettage, electrodessication, cryotherapy, and PDT

Curettage and electrodessication may be considered for small and low-risk primary SCC, according to NCCN guidelines. As indicated by NCCN guidelines, cryotherapy could be a treatment option in selected cases of low-risk SCC, while there is scarce evidence regarding efficacy of PDT for invasive SCC and it should not be performed in these cases.

Figure 4. Main therapeutic indications for locally advanced and metastatic cutaneous SCC as indicated in the “European interdisciplinary guideline on invasive squamous cell carcinoma of the skin: Part 2” in 2020. SCC, cutaneous squamous cell carcinoma; RT, radiotherapy; EGFRi, epidermal growth factor receptor inhibitors.
Surgery for regional nodal disease

Patients affected by SCC nodal metastases should be surgically treated similarly to patients affected by melanoma or Merkel cell carcinoma. If surgery is not possible due to patient-related factors, a non-surgical approach such as immunotherapy with cemiplimab should be considered by the MWG. The regional lymph node dissection is the surgical treatment of choice in case of nodal metastasis.11,59,60

Considering patients affected by SCC and negative lymph node, an elective or prophylactic lymph node dissection is not recommended due to the low rate of nodal metastases, the high morbidity and the limited evidence in patients with mucosal head and neck SCC.61,62

Electrochemotherapy

ECT is an alternative treatment for unresectable SCC consisting of an intravenous injection of a chemotherapy agent, such as cisplatin or bleomycin, combined with local electric pulses that permeabilize tumor cell membranes to increase its cytotoxicity.63

In some retrospective studies and one meta-analysis, it has been reported that 20-70% of patients treated with ECT presented a good local response and disease control, while, in a prospective study EURECA on SCC patients the rate of complete response at 2-months follow-up was 55% with 4% rate of progression only.64-67

Radiation

RT could be an alternative treatment in case of patients affected by SCC who are not eligible for surgery such as locally advanced tumor, multi-morbidity or frail elderly patient at high risk for surgery, or patient that refuse surgery. Otherwise, surgery must be chosen wherever possible because RT presents lower cure rates and many cases of aggressive post-treatment recurrence have been observed. In a meta-analysis of 14 observational studies, a 6.4% average rate of local tumor recurrence after the first-line RT in 1018 primary SCC has been reported.68

RT could also be considered as an esthetic option for SCC localized on neck and head or as a functional option for SCC localized on sensitive areas such as lips or eyelids. Furthermore, radical primary RT could be used for small SCC.69,70

AEs related to RT are rare and consist of radiodermatitis, hypo/hyperpigmentation, and telangiectasia. Furthermore, RT is not recommended in patients with genetic disorders such as Gorlin syndrome or ataxia telangiectasia and others for the higher risk of radiosensitivity.71

Systemic treatment

Most advanced SCC, including locally advanced or metastatic lesions, may be non-resectable. Locally advanced SCC has been defined as a non-metastatic SCC, not amenable to either surgery or RT with reasonable hope for cure, because of multiple recurrences, large extension, bone erosion or invasion, or deep infiltration beyond subcutaneous tissue into muscle or along nerves, or else tumors in which curative resection would result in unacceptable complications, morbidity or deformity.11

Metastatic SCC includes loco-regional metastatic SCC with in-transit metastases or metastasis to regional lymph nodes, or distant metastatic SCC.11 Until recently, treatment options have been off-label chemotherapy or anti-epidermal growth factor receptor (EGFR) therapies. However, the clinical evidence for these options is limited and chemotherapy is associated with a high risk of significant adverse events, especially in older patients.72

The therapy with immune checkpoint inhibitors and targeted therapy could be an alternative treatment in many cases of aggressive post-treatment recurrences, large extension, bone erosion or invasion, or deep infiltration beyond subcutaneous tissue into muscle or along nerves, or else tumors in which curative resection would result in unacceptable complications, morbidity or deformity.11

Table 3. TNM Staging Classification for Cutaneous Carcinoma of the Head and Neck according to the American Joint Committee on Cancer (AJCC) (8th ed., 2017). Definitions for T and clinical N.

<table>
<thead>
<tr>
<th>T</th>
<th>Primary Tumor</th>
</tr>
</thead>
<tbody>
<tr>
<td>TX</td>
<td>Primary tumor cannot be assessed</td>
</tr>
<tr>
<td>Tis</td>
<td>Carcinoma in situ</td>
</tr>
<tr>
<td>T1</td>
<td>Tumor smaller than or equal to 2 cm in greatest dimension</td>
</tr>
<tr>
<td>T2</td>
<td>Tumor larger than 2 cm, but smaller than or equal to 4 cm in greatest dimension</td>
</tr>
<tr>
<td>T3</td>
<td>Tumor larger than 4 cm in maximum dimension or minor bone erosion or perineural invasion or deep invasion*</td>
</tr>
<tr>
<td>T4</td>
<td>Tumor with gross cortical bone/marrow, skull base invasion and/or skull base foramen invasion</td>
</tr>
<tr>
<td>T4a</td>
<td>Tumor with gross cortical bone/marrow invasion</td>
</tr>
<tr>
<td>T4b</td>
<td>Tumor with skull base invasion and/or skull base foramen involvement</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>cN</th>
<th>Regional Lymph Nodes</th>
</tr>
</thead>
<tbody>
<tr>
<td>NX</td>
<td>Regional lymph nodes cannot be assessed</td>
</tr>
<tr>
<td>N0</td>
<td>No regional lymph node metastasis</td>
</tr>
<tr>
<td>N1</td>
<td>Metastasis in a single ipsilateral lymph node, 3 cm or smaller in greatest dimension and ENE(−)</td>
</tr>
<tr>
<td>N2</td>
<td>Metastasis in a single ipsilateral node larger than 3 cm but not larger than 6 cm in greatest dimension and ENE(−); or metastases in multiple ipsilateral lymph nodes, none larger than 6 cm in greatest dimension and ENE(−); or in bilateral or contralateral lymph nodes, none larger than 6 cm in greatest dimension and ENE(−)</td>
</tr>
<tr>
<td>N2a</td>
<td>Metastasis in a single ipsilateral node larger than 3 cm but not larger than 6 cm in greatest dimension and ENE(−)</td>
</tr>
<tr>
<td>N2b</td>
<td>Metastases in multiple ipsilateral nodes, none larger than 6 cm in greatest dimension and ENE(−)</td>
</tr>
<tr>
<td>N2c</td>
<td>Metastases in bilateral or contralateral lymph nodes, none larger than 6 cm in greatest dimension and ENE(−)</td>
</tr>
<tr>
<td>N3</td>
<td>Metastasis in a lymph node larger than 6 cm in greatest dimension and ENE(−); or metastasis in any node(s) and clinically overt ENE [ENE(+) ]</td>
</tr>
<tr>
<td>N3a</td>
<td>Metastasis in a lymph node larger than 6 cm in greatest dimension and ENE(−)</td>
</tr>
<tr>
<td>N3b</td>
<td>Metastasis in any node(s) and ENE(+)</td>
</tr>
</tbody>
</table>

*Deep invasion is defined as invasion beyond the subcutaneous fat or > 6 mm (as measured from the granular layer of adjacent normal epidermis to the base of the tumor); perineural invasion for T3 classification is defined as tumor cells within the nerve sheath of a nerve lying deeper than the dermis or measuring 6.3 mm or larger in caliber; or presenting with clinical or radiographic involvement of named nerves without skull base invasion or transgression. Note: A designation of “U” or “L” may be used for any N category to indicate metastasis above the lower border of the cricoid (U) or below the lower border of the cricoid (L). Similarly, clinical and pathological extranodal extension (ENE) should be recorded as ENE(−) or ENE(+) .
Other inhibitors of PD-1/PDL-1-axis are under investigation and Pembrolizumab, a PD-1 inhibitor, has been recently approved by FDA on the basis of a phase II trial for patients with recurrent or metastatic cutaneous SCC that is not curable by surgery or radiation.\textsuperscript{77}

The safety and the efficacy of intravenous cemiplimab was investigated in a phase I study for expansion cohorts of patients with locally advanced or metastatic cutaneous squamous-cell carcinoma (NCT02383212), as well as in a pivotal phase II study for a cohort of patients with metastatic disease (metastatic-disease cohort or group 1) (NCT02760498 EMPOWER-CSCC 1).\textsuperscript{75} Patients who had undergone organ transplantation and patients with hematologic malignancies or undergone organ transplantation and patients with hematologic malignancies or any immunosuppressive conditions were excluded in both studies. The patients were treated with cemiplimab (3 mg per kilogram of body weight) intravenously every 2 weeks for up to 48 weeks (phase I) or 96 weeks (phase II) unless stopped due to disease progression or non-tolerable toxic effects. Patients were assessed for a response every 8 weeks. The primary outcomes of the phase I study were the safety and adverse event profile, while the response rate assessed by independent central review was the primary outcome of the phase II study. In the phase I study, a response to cemiplimab was observed in 13 of 26 patients (50%; 95% CI = 30 to 70). In the metastatic-disease cohort of the phase II study, a response was observed in 28 of 59 patients (47%; 95% CI = 34 to 61). The median follow-up was 11.1 months in the phase I and 7.9 months in the metastatic-disease cohort of the phase II study. Among the 28 patients who had a response, the duration of response exceeded 6 months in 57%, and 82% continued to have a response and to receive cemiplimab at the time of data cutoff. The treatment was generally well tolerated with only 7% of patients stopping therapy due to adverse events. The most commonly reported adverse events (15%) usually occur with immune checkpoint inhibitors, such as diarrhea, fatigue, nausea, constipation and rash. In 3 of 11 patients in the phase II cohort who died during the study, death was associated with a non-treatment emergent adverse event. Based on this study, cemiplimab was

\begin{table}[h]
\centering
\begin{tabular}{|c|c|}
\hline
\textbf{pN} & \textbf{Regional Lymph Nodes} \\
\hline
NX & Regional lymph nodes cannot be assessed \\
N0 & No regional lymph node metastasis \\
N1 & Metastasis in a single ipsilateral lymph node, 3 cm or smaller in greatest dimension and ENE(–) \\
N2 & Metastasis in a single ipsilateral lymph node, 3 cm or smaller in greatest dimension and ENE(+) or larger than 3 cm but not larger than 6 cm in greatest dimension and ENE(–); or metastases in multiple ipsilateral lymph node(s), none larger than 6 cm in greatest dimension and ENE(–); or in bilateral or contralateral lymph node(s), none larger than 6 cm in greatest dimension, ENE(–) \\
N2a & Metastasis in single ipsilateral node 3 cm or smaller in greatest dimension and ENE(+) or a single ipsilateral node larger than 3 cm but not larger than 6 cm in greatest dimension and ENE(–) \\
N2b & Metastases in multiple ipsilateral nodes, none larger than 6 cm in greatest dimension and ENE(–) \\
N2c & Metastases in bilateral or contralateral lymph node(s), none larger than 6 cm in greatest dimension and ENE(–) \\
N3 & Metastasis in a lymph node larger than 6 cm in greatest dimension and ENE(–); or in a single ipsilateral node larger than 3 cm in greatest dimension and ENE(+) or multiple ipsilateral, contralateral, or bilateral nodes, any with ENE(+); or a single contralateral node of any size and ENE(+) \\
N3a & Metastasis in a lymph node larger than 6 cm in greatest dimension and ENE(–) \\
N3b & Metastasis in a single ipsilateral node larger than 3 cm in greatest dimension and ENE(+) or multiple ipsilateral, contralateral, or bilateral nodes, any with ENE(+); or a single contralateral node of any size and ENE(+) \\
M & Distant Metastasis \\
M0 & No distant metastasis \\
M1 & Distant metastasis \\
G & Histologic Grade \\
GX & Grade cannot be assessed \\
G1 & Well differentiated \\
G2 & Moderately differentiated \\
G3 & Poorly differentiated \\
G4 & Undifferentiated \\
\hline
\end{tabular}
\caption{TNM Staging Classification for Cutaneous Carcinoma of the Head and Neck according to the American Joint Committee on Cancer (AJCC) (8th ed., 2017). Definitions for Pathological N and metastasis.}
\end{table}
approved in September 2018 by the FDA. In 2020, Migden et al. 75 reported the clinical activity of cemiplimab from the primary analysis of patients with locally advanced cutaneous SCC (group 2) from the pivotal phase II study (NCT02760498). Between 2016 and 2018, 78 patients were enrolled and treated with cemiplimab (3 mg/kg) intravenously over 30 min every 2 weeks for up to 96 weeks. Tumor measurements were done every 8 weeks. An objective response was observed in 34 (44%; 95% CI 32-55) of 78 patients. The best overall response was ten (13%) patients with a complete response and 24 (31%) with a partial response. Grade 3-4 treatment-emergent adverse events occurred in 34 (44%) of 78 patients. The most common were hypertension in six (8%) patients and pneumonia in four (5%). Serious treatment-emergent adverse events occurred in 23 (29%) of 78 patients. One treatment-related death was reported that occurred after onset of aspiration pneumonia. Thus, cemiplimab showed antitumor activity and an acceptable safety profile in patients with locally advanced cutaneous squamous cell carcinoma for whom there was no widely accepted standard of care. In the meantime, Rischin et al. 76 published outcomes of the primary analysis of fixed dose cemiplimab 350 mg intravenously treatment every 3 weeks (Group 3) and provide a long-term follow-up after the primary analysis of weight-based cemiplimab 3 mg/kg intravenously every 2 weeks (Q2W) (Group 1) among metastatic SCC patients in the pivotal study phase II. For Group 3 (n=56) and Group 1 (n=59), median follow-up was 8.1 and 16.5 months, respectively. ORR per ICR was 41.1% (95% CI, 28.1% to 55.0%) in Group 3, 49.2% (95% CI, 35.9% to 62.5%) in Group 1, and 45.2% (95% CI, 35.9% to 54.8%) in both groups combined. Per ICR, Kaplan-Meier estimate for DOR at 8 months was 95.0% (95% CI, 69.5% to 99.3%) in responding patients in Group 3, and at 12 months was 88.9% (95% CI, 69.3% to 96.3%) in responding patients in Group 1. Per INN, ORR was 51.8% (95% CI, 38.0% to 65.3%) in Group 3, 49.2% (95% CI, 35.9% to 62.5%) in Group 1, and 50.4% (95% CI, 41.0% to 59.9%) in both groups combined. Overall, the most common adverse events regardless of attribution were fatigue (27.0%) and diarrhea (23.5%). Therefore, durable responses and similar safety profile have been observed in both weight-based and fixed-dosing groups.76 A further update of this trial has been presented at 10th World Congress of Melanoma in 2021 with a prolonged follow-up at 43 months. Compared to previous analyses, the 43-month follow-up data demonstrated incremental improvements in DOR with cemiplimab treatment across all locally advanced SCC study groups, as well as improvements in ORR and complete response rate on the cemiplimab 350 mg every 3 weeks regimen. Furthermore, there were no new safety signals compared with previous reports on cemiplimab in advanced cutaneous SCC.

Staging

TNM staging classification for head and neck BCC and cutaneous SCC and prognostic stage groups according to the AJCC 8th edition is represented in Tables 3-5.

Follow up

The patient should be regularly monitored to recognize any new NMSC or relapses of the tumor and, mainly for locally advanced or metastatic cases, to manage the HPI therapy for BCC or immunotherapy for SCC. Furthermore, it is essential to educate the patient regarding the correct photoprotection and the periodic self-control of skin lesions.

Basal cell carcinoma follow-up

Considering that a patient with a previous diagnosis of BCC has a 15% higher risk to develop another BCC in one year and 35% in 5 years and that the risk increases in case of multiple previous BCCs, it is important to advice a periodic follow-up to the patient. This follow-up should be performed by dermatologists and should include the inspection of the entire body. Monitoring in the first two years is essential. Furthermore, dermatological examination should be performed every 6-12 months (Table 6).

Squamous cell carcinoma follow-up

Considering that 95% of relapses and the same percentage of metastases occur within the first five years from SCC diagnosis and that 30-50% of patients may present a second SCC within 5 years, patients affected by SCC requires a long-term follow-up.

SCC patients should be monitored with regular physical exams that include complete skin and regional lymph node examination. The frequency of follow-up depends on the risk of SCC and it is indicated in Table 6.

Conclusions

The ICP for NMSC represents an innovative strategy to support the highest quality health care system, favouring all necessary procedures for the patients, optimizing the necessary timing, and guaranteeing an updated clinical knowledge of the various health professionals involved. The hub-and-spoke model is essential for the organisa-

Table 5. Prognostic stage groups according to the American Joint Committee on Cancer (AJCC) (8th ed., 2017).

<table>
<thead>
<tr>
<th>Stage</th>
<th>T</th>
<th>N</th>
<th>M</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>Tis</td>
<td>N0</td>
<td>M0</td>
</tr>
<tr>
<td>I</td>
<td>T1</td>
<td>N0</td>
<td>M0</td>
</tr>
<tr>
<td>II</td>
<td>T2</td>
<td>N0</td>
<td>M0</td>
</tr>
<tr>
<td>III</td>
<td>T3</td>
<td>N0</td>
<td>M0</td>
</tr>
<tr>
<td></td>
<td>T1</td>
<td>N1</td>
<td>M0</td>
</tr>
<tr>
<td></td>
<td>T2</td>
<td>N1</td>
<td>M0</td>
</tr>
<tr>
<td></td>
<td>T3</td>
<td>N1</td>
<td>M0</td>
</tr>
<tr>
<td>IV</td>
<td>T1</td>
<td>N2</td>
<td>M0</td>
</tr>
<tr>
<td></td>
<td>T2</td>
<td>N2</td>
<td>M0</td>
</tr>
<tr>
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<td>N2</td>
<td>M0</td>
</tr>
<tr>
<td></td>
<td>Any T</td>
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<td>M0</td>
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<td></td>
<td>T4</td>
<td>Any N</td>
<td>M0</td>
</tr>
<tr>
<td></td>
<td>Any T</td>
<td>Any N</td>
<td>M1</td>
</tr>
</tbody>
</table>

Table 6. BCC and SCC recommended follow-up (NCCN BCC and SCC version 2021).

<table>
<thead>
<tr>
<th>BCC</th>
<th>6-12 months</th>
</tr>
</thead>
<tbody>
<tr>
<td>Low-risk SCC</td>
<td>Every 3-12 months for 2 years, then every 6-12 months for 3 years, then annually for life</td>
</tr>
<tr>
<td>High-risk SCC</td>
<td>Every 3-6 months for 2 years, then every 6-12 months for 3 years, then annually for life</td>
</tr>
<tr>
<td>Very high-risk SCC</td>
<td>Every 3-6 months for 2 years, then every 6 months for 3 years, then every 6-12 months for life</td>
</tr>
</tbody>
</table>

[Dermatology Reports 2021; 13:9278]
tion of different health care structures to guarantee the best management and treatment for patients affected by NMSC. Considering the increasing incidence of these very common tumors and the high impact on the sanitary system, it is crucial to create an efficient and dedicated ICP with a valid organization within the different geographical areas.

References

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