Management of superficial basal cell carcinoma: focus on imiquimod

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Abstract: Superficial basal cell carcinoma comprise up to 25% of all histological sub-types. They are more likely to occur on younger persons and females and although generally more common on the trunk, also occur frequently on the exposed areas of the head and neck especially in areas of high sun exposure. In the last decade, new treatment options such as topical applications that modify the immune response have been trialed for effectiveness in treating these lesions. Imiquimod 5% cream has been shown to stimulate the innate and cell mediated immune system. The short-term success of imiquimod 5% cream in randomized controlled trials comparing different treatment regimes and dosing as a treatment for small superficial basal cell carcinoma (BCC) not on the face or neck is in the range of 82% for 5 times per week application. A high proportion of participants with good response rates to topical treatment (58%–92%) experience local side effects such as itching and burning, less commonly erosion and ulceration, but the proportion of participants ceasing treatment has not been high. To date one long-term study indicates a treatment success rate of 78%–81% and that initial response is a predictor of long-term outcome. Recurrences tend to occur within the first year after treatment. Future research will compare this preparation to the gold standard treatment for superficial BCC – surgical excision.

Keywords: superficial basal cell carcinoma, imiquimod, skin cancer

Introduction
Basal cell carcinoma (BCC) is the most common skin cancer in fair skinned populations and the incidence of BCC continues to rise by approximately 10% per year. BCC can be classified into different histological sub-types according to their growth pattern.1,2 The different histological sub-types broadly distinguish between nodular, superficial and infiltrative and other less frequently occurring types according to their growth patterns.3 Micro-nodular and infiltrative sub-types are recognized as high risk and should rarely be treated other than surgically. Apart from the histological appearance, other well-recognized factors that can influence the prognosis and outcomes for treated BCC are the size and site of the tumor, how clearly the margins are defined and failure of previous treatment.2

The aim of this review is to examine the literature to identify the factors that can help a clinician decide whether or not the topical application imiquimod 5% cream is an appropriate treatment for a patient with the superficial sub-type of BCC (sBCC). The paper will outline the evidence supporting its use and the situations in which it is most likely to be successful, the most appropriate treatment regime and the expected disadvantages and side-effects of this treatment modality for this type of lesion.
Review criteria
The information for this review was compiled by searching the PubMed, Cochrane, MD Consult and DARE databases and hard copy and electronic journals of dermatology, cancer and therapeutics. The search terms used included ‘reviews’ and ‘trials’, ‘randomized controlled trials’ ‘basal cell carcinoma’ ‘superficial basal cell carcinoma, ‘sBCC’ and ‘BCC’ and ‘treatment with imiquimod’ ‘cost of treatment.’ Electronic early-release publications were also included. Only articles published in English were considered. When possible, primary sources have been quoted. Full articles were obtained and references were checked for additional material when appropriate. References were chosen on the basis of the best clinical or laboratory evidence, especially if the work had been corroborated by published work from other centers.

Definition and epidemiology of sBCC
The superficial type BCC makes up 25%-26% of all BCCs in sun-exposed Australians with age-standardized incidence rates of 337/100,000/year for males and 251/100,000/year for females compared with 15%-16% of the total types in Europe. From reports in Europe the superficial type is more likely to occur on the trunk and in the younger age groups especially in females. In Australia, although still less common than the nodular type, compared with Europe, there are proportionately more sBCC, and in females the incidence is maximum in the 40-49 years age group. The relative tumor density is highest on the face although the highest proportion of sBCC is on the trunk in males. Clinically, sBCC are erythematous, slightly scaly, well-defined patches, most commonly found on the trunk and the limbs. However, as mentioned in highly sun-exposed population are also common on the face. They grow slowly and can be confused with patches of psoriasis or eczema or even fungal infections. Although the lesion is more likely to be superficial, they can be large (>10 mm) identifying the margin when excising the lesions can be quite difficult, and may be inadequate excised and so recur. Recurrence of excised BCC can occur up to 10-20 years post treatment and may potentially exceed 50%. However, this does not necessarily apply to sBCC types. Understanding the etiology and histopathology of these lesions is important because it helps to explain the difficulty in clinical diagnosis and therefore the role of an immune response modifier in treatment.

Etiology
Evidence suggests that BCC may arise from the pluripotent cells in the basal layer or follicles of the skin. Mutations in the PATCHED (PTCH) gene, a negative regulator of hedgehog signaling, located on chromosome 9 appear to be responsible for the tumor growth in basal cell nevus syndrome (BCNS), an autosomal condition characterized by multiple BCCs. Mutations in the PTCH gene have also been reported in sporadic cases of BCC and xeroderma pigmentosum associated with BCC. A recent study that focused on cases of sporadic BCC showed that almost half of the tumors examined bore PTCH gene mutations, and the percentage and type of these mutations did not differ significantly among patient groups that had regular, multiple, or early-onset BCC.

In general, BCCs are made up of nests of cells representing the basal epidermal layers with peripherally palisading cells. The cells have hyperchromatic nuclei and scant cytoplasm. The nests of tumor cells are surrounded by stroma. Intercellular bridges are not visible on light microscopy. Mitotic figures are common but the whole appearance is one of uniformity rather than anaplasia. Ulceration is not uncommon in large tumors. In long-standing or aggressive lesions, extension is often diffuse or in the paths of cutaneous adnexae. Perineural invasion is seen in about 1% of cases, more frequently in aggressive forms of BCC. A variable inflammatory infiltrate is often present, usually with a majority of (CD4+) T cells. There is a prominent stroma arranged in bundles around the tumor masses. The (multifocal) superficial type is characterized by numerous small nests of tumor cells usually attached to the undersurface of the epidermis by a broad base. Approximately 10%-15% of all BCCs are of this type. This is the most common pattern seen in BCCs of the shoulder.

Treatment options for sBCC
There are a variety of treatment options for BCC as a whole, including the superficial sub-type; however, little research has been done comparing the treatment options for this specific sub-type taking into account such factors as age of the patients, body site of the lesions, operator skills, with adequate follow-up and assessing in addition to recurrence rates, the patient outcomes of cosmesis and quality of life (QOL).

In examining the role of topical applications for the sBCC it may therefore be necessary to define more clearly where surgery is inappropriate or not first line treatment so that the less invasive options such as topical applications can be recommended.

Surgery and destructive modalities in the treatment of sBCC
Included in the surgical option for treatment are a number of different modalities including complete primary excision,
Mohs surgery and the destructive treatments of curettage and cautery, and cryosurgery. These procedures have pros and cons when it comes to sBCC.

The overall treatment with the best evidence base for primary sBCC and other subtypes is still surgical with recurrence rates of <2% and good cosmetic results. This outcome depends significantly on the size of the lesion. For a tumor ≤2 cm, the control rate at 5 years is expected at 95% with a tumor ≤5 cm and deeply invading beyond subcutaneous tissues having a control rate from as low as 50%. The reason for a good outcome for surgery is the ability to examine the histopathology of specimens and to determine the adequacy of margins. Successful primary excision reduces the recurrence of any type of BCC including sBCC. In general, nodular and superficial sub-types, which account for the majority of lesions, do not have aggressive features and have a higher rate of complete excision and lower recurrence rates. Nevertheless, up to 38% of tumors in one series were mixed sub-types of BCC, and so what appears to be a simple sBCC may behave like the most aggressive sub-component which is unrecognized clinically and may be missed even when biopsied. GPs in Australia who treat most sBCC in their practices report treating over half (57%) with surgery.

Mohs surgery in general would not be undertaken for a sBCC unless a recurrent lesion on a high-risk area. It is a highly specialized surgical procedure combining staged procedures with examination of all margins, and aims to remove all tumor, with tissue sparing and reconstruction if necessary. It is a procedure not available in all areas of high skin cancer incidence, and it is generally reserved for high-risk or recurrent lesions and where there is perineural or perivascular involvement.

The strength of the evidence for use of curettage and cautery alone is weak and limited to evidence provided by multiple time series and successful results depend on careful selection of low-risk lesions, preferably not on the face where it is less successful and on the skill and experience of the operator. It has been a preferred method used by dermatologists to treat primary nBCC and sBCC < 1.5 cm although recent data on recurrence rates are limited. It has been used in conjunction with newer treatments such as topical applications and PDT especially for nodular BCC and will be discussed further below in association with the use of imiquimod cream.

Cryosurgery is a common form of treatment for BCC in dermatological practice where it is most suitable for primary, well-defined superficial lesions of low risk on other than the head and neck. Many large older studies demonstrate satisfactory cure rates. One review including studies up to and including 1997, suggested recurrence rates with this modality of from 4% to 17%. A more recent review revealed that there is a dearth of prospective, randomized controlled trials with a 5-year follow-up and that recurrence rates are variable. Success is technique dependent and one review of multiple series reported a recurrence rate of 7.5% compared with other treatment modalities. This technique has been successfully used to treat sBCC with single freeze thaw cycles achieving cure rates of 96%. One of the disadvantages of this type of treatment is the perioperative pain, tenderness, blistering and sloughing of necrotic tissue, with possible hypopigmentation and scarring as the outcome and the difficulty associated with identifying recurrence in scar tissue.

In the most recent Cochrane review of interventions for BCC only three randomized controlled trials (RCTs) were identified comparing surgical excision with an alternative treatment. First, surgical excision versus radiotherapy (RT) which does not specifically differentiate between types of BCC although sBCC were included. Second, surgical excision versus Mohs which looks at high-risk BCCs only, and third, surgical excision versus photodynamic therapy (PDT) which includes only nodular BCC. Cryosurgery has been compared with radiotherapy and excision and PDT. Although the trial where cryosurgery was compared with PDT identifies sBCC (n = 39) in treatment it does not mention how these specifically fared following the intervention compared with other types. The latter trial of 5-year follow-up of treatment of sBCC by cryosurgery (105 lesions on 58 patients) versus PDT looked at (n = 114 lesions) of small size on 60 patients and although the outcome significantly favored PDT for patient tolerability and cosmesis, there was still a high recurrence rate of 20% for cryosurgery compared with 22% with PDT. These trials are outlined in Table 1. No randomized controlled trials (RCTs) are reported comparing cautery and curettage with any other treatment modality although curettage was used to debulk tumors in one study.

Photodynamic therapy (PDT)

This modality uses topical photosensitizing agents aminolevulinic acid (ALA) and MAL, its methylated ester, which are applied topically to the lesion and allowed to stay on the skin and be absorbed for a period of time before being subjected to a specific light source. These agents are relatively selectively concentrated in the lesion and when exposed to the light source, usually in the wavelength of 620 to 670 nm, this concentration results in photodestruction of the lesion. Other light sources are also used in PDT including lasers, filtered xenon arc and metal halide lamps, fluorescent lamps and light-emitting diodes. Studies have suggested there is a place for the treatment of
sBCC with PDT, although they may not clear with a single treatment and that results are better with two cycles. One RCT examined sBCC (n = 245 lesions on 83 patients) using topical ALA and 2 different light sources and there was no significant difference in the clinical or cosmetic outcomes at 6 months. As mentioned above, PDT has been compared with cryosurgery in an RCT, but not specifically for sBCC and not against other noninvasive modalities such as topical 5-fluorouracil (5FU) which has Federal Drug Authority approval for treatment of sBCC where conventional methods are impractical, or the immune-response modifier imiquimod.

### Topical applications
A body of literature and trials are now available looking at topical applications in the treatment of sBCC, although it is not always easy to determine whether the sub-type of BCC has been clearly defined. In addition, a successful outcome for treatment needs to be explicit and it is recommended that 5- to 10-year local control rates or recurrence rates are legitimate measures although most studies do not go beyond 5 years. As with any treatment modality the desired primary outcome is to totally eradicate the tumor and secondly to do so with as good a cosmetic outcome and patient acceptance as possible yet comparative data for different treatments is sparse, with most topical application studies now either comparing different treatment regimes or long-term follow-up rather than against other intervention types.

For example, the actual clearance rate for treatment for sBCC with 5FU is currently not known because of a lack of adequate trials. The only open-label randomized trial available includes superficial and nodular BCC and compared six different treatment regimes. Overall there was an 8% treatment failure for sBCC when followed up by histological examination at 3 months. One 1-year follow-up study comparing 5FU with imiquimod for actinic keratoses (AK) is reported.

### Focus on imiquimod for sBCC

#### Mode of action
Imiquimod is a synthetic preparation that acts as an immune response modifier and its clinical effect comes from cytokine activation of the immune system and possibly also from induced apoptosis in BCC cells. It has been used now for some years in the treatment of genital warts as a topical application where it stimulates the production of cytokines including interferon alpha through activation of Toll-like receptors (TLR) -7 on antigen presenting cells. It promotes

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**Table 1** Comparison trials of treatment where sBCC are specifically included

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<th>Intervention trial</th>
<th>Inclusion</th>
<th>Follow up</th>
<th>Outcome</th>
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<tr>
<td>Excision vs RT</td>
<td>Included primary tumors sBCC (n = 36) among other types, face, &lt;40 mm</td>
<td>Follow-up 4 years: histopathology, primary and secondary</td>
<td>Primary: recurrence and secondary cosmesis both favored surgery</td>
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<td>Favored surgery for cosmesis, and no significant difference for clinical recurrence</td>
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<td>Excision vs cryosurgery</td>
<td>n = 103, sBCC, nBCC head and neck &lt;2 cm</td>
<td>Follow-up 1 year: clinical superficial (cryosurgery n = 8, excision n = 6)</td>
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<td>Note: curettage also used</td>
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<td>nodular cryosurgery n = 40, excision n = 42</td>
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<td>Surgical excision vs crye</td>
<td>n = 100, sBCC n = 4 (all treated with excision)</td>
<td>Follow-up 5 years n = 85</td>
<td>Statistically non-significant (C&amp;C 19.6% vs SE 8.4%)</td>
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<td>Cryosurgery vs RT</td>
<td>? histological sub-types n = 93</td>
<td>Follow-up 1 year: histopathology</td>
<td>Favored RT comparable cosmetic</td>
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<tr>
<td>Cryosurgery vs PDT</td>
<td>sBCC n = 39, nod n = 49</td>
<td>Follow-up 1 year: Histopathology (n = 83)</td>
<td>Significandy favored PDT for patient tolerability and cosmesis. Recurrence 25% (11 of 44) for ALA-PDT and 15% (6 of 39) for cryosurgery.</td>
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<tr>
<td>Cryosurgery vs PDT</td>
<td>sBCC n = 245, thickness &lt;1 mm, diam &lt;3 mm</td>
<td>Follow-up 5 years (n = 193)</td>
<td>Significandy favored PDT for patient tolerability and cosmesis, but no sig diff in recurrence (20% cryo vs 22% PDT)</td>
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<td>Curettage and cautery</td>
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**Abbreviations:** ALA, aminolevulanic acid; RT, radiotherapy; PDT, photodynamic therapy; C&C, curettage and cryosurgery; SE, surgical excision; sBCC, superficial basal cell carcinoma.
the innate immune response and the cell-mediated immune pathway. This immune modification has also been shown to produce anti-proliferative and anti-tumor activity.\(^{36,37}\)

In examining biopsies of sBCC before and after treatment and undertaking gene studies and immunohistochemistry, it has been found that imiquimod has an anti-tumor effect on sBCC. This effect is mediated by the stimulation of the dendritic cells in the epidermis and dermis to release interferon (IFN) \(\alpha\) and other cytokines, thus activating the innate immune system with activity against tumor cells resulting in apoptosis.\(^{38,39}\) A recent double blind placebo-controlled randomized parallel group trial designed to identify the early immunological events caused by the application of imiquimod when used to treat BCC (type not indicated) found that the early mode of action was characterized by the stimulation of a large number of genes (637) which were highly specific to imiquimod action, 98 of which confirmed previous involvement of interferon \(\alpha\) (IFN-\(\alpha\)). With the remaining genes IFN \(\gamma\) transcription was more prevalent than the former. The most striking effects after several days of application were on cytotoxic mechanisms with significant increases and induction of recruitment and activation of T and natural killer (NK) cells within the BCC. It was hypothesized that that it is these secondary immune effect mechanisms that induce destruction of target cells. It appears therefore that imiquimod links multiple immune pathways and of these IFN-\(\alpha\) plays a consistent but not exclusive role.\(^{40}\)

A further possible action of imiquimod was demonstrated when it was applied in vitro to squamous cell carcinomas (SCC). It appeared to neutralize the failure of the tumors to express vascular E-selectin. Through this it was able to allow the skin-homing T cells access to the tumor to promote its destruction. In addition, it reduced the number of tumor infiltrating T regulatory cells thus tipping the balance of T cells within the tumor towards CD8+ T cells. Furthermore, it induced interleukin 6 (IL-6) production from the effector T cells. These actions may also be relevant to sBCC.\(^{41}\)

It appears to have minimal impact on normal skin, does not increase ultraviolet (UV) induced change to epidermal keratinocytes or DNA and the systemic absorption is minimal, with <0.9% of the dose excreted in urine and <0.2% excreted in feces, although flu-like symptoms and headache can occur while using the cream especially in older patients.\(^{42,43}\)

The effect of imiquimod has also been studied by a new technique of confocal microscopy. This is a non-invasive optical technique that can be used to evaluate skin diseases without biopsy. Unfortunately it is very expensive equipment that would not be available in all centers. Nevertheless in a small case control study and in one case report the investigators were able to demonstrate the local immune response following therapy with imiquimod and demonstrated how the previously diseased skin improved over time. Histology of the tissue was also carried out to confirm findings.\(^{44,45}\)

**Uses**

In 2004, imiquimod (Aldara\(^{®}\); 3M, St. Paul, MN) was approved by the United States Food and Drug Administration (FDA) for the topical treatment of sBCC, based on two double-blind, vehicle-controlled clinical studies.\(^{46,47}\) The indications for use were for the topical treatment of biopsy-confirmed, primary superficial BCC in immunocompetent adults, with a maximum tumor diameter of 2.0 cm, located on the trunk (excluding anogenital skin), neck, or extremities (excluding hands and feet), only when surgical methods are medically less appropriate and patient follow-up can be reasonably assured.\(^{48}\)

In Australia, following results of the above Phase III trials where clearance rates were 82%, imiquimod cream 5% is approved for use by the Therapeutic Goods Administration for biopsy-proven primary sBCC, and solar keratoses. Dosing for the former is application to the lesion and 5 mm surrounding skin 5 times per week for 6 weeks. Treatment breaks are allowed for severe inflammatory reactions.\(^{11,47}\) For subsidized use under the Pharmaceutical Benefits Scheme it is approved for the “Treatment of superficial basal cell carcinoma (BCC) in immunocompetent patients who cannot have surgical excision, cryotherapy, or curettage with diathermy. The lesion must be previously untreated and the diagnosis confirmed by biopsy, and the patient or carer must be able to follow the dosing regimen.”\(^{49}\)

Approval for use in the UK is for treatment of superficial BCCs that measure up to 2 cm across (diameter) on the trunk, neck, arms and legs (including hands and feet).\(^{50}\)

**Trials: what do they tell us?**

Initial trials of imiquimod 5% cream for the treatment of skin cancer focused on sBCC, and were drug-company sponsored studies in prospective open-label dose-response studies in multiple centers in Australia, Europe and the United States. The inclusion criteria for these studies were primary histologically confirmed sBCC lesions, not previously treated, of <2 cm diameter or 0.5 mm\(^2\) and in a position where it could be later excised. The patients were to have no skin conditions, and be immunocompetent. These first trials indicated that imiquimod could be an effective
treatment for sBCC and that an appropriate treatment regime would be a maximum of a daily application, higher dosing causing more and intolerable side effects. The length of treatment was also tested in these studies with tumor treatment of varying times from 6 to 12 weeks, as was the ability to determine clinically whether the tumor had been cleared.31,52 The predictive value of clinical diagnosis of tumor non-clearance was 39% by the investigators, but when clinical diagnosis of positive clearance was compared with histology results investigators were correct in 100% of cases.46 The results of these trials suggested that a clearance rate, based on clinical and histological examination, could be in the range of 73% to 88% with a maximum clearance rate with daily applications for 6 weeks.

Of the nine RCTs reported on imiquimod use in the treatment of basal cell carcinoma in the most recent Cochrane review,10 only four relate to sBCC, with 128 patients, 724 patients 166 patients and 35 patients respectively35,46,47,51 with another relating to sBCC that was a randomized open label study with 93 patients with sBCC.54 These studies are outlined in Table 2. A 3-year primary outcome with 5-year follow-up RCT comparing the use of imiquimod with excisional surgery for the treatment of nodular and sBCC is underway in the United Kingdom.54 All the studies have compared the treatment intervention with vehicle and with different doses and dosing regimes. In the most recent Cochrane review all except the study undertaken by Sterry et al were determined to be of medium quality.

In summary, for the larger trials identified, the clearance rates outlined in Table 2 for once-daily 5 to 7 times per week dosing, ranged from overall (clinical and histological combined) 73% to 75%,47 77%43 to 81% to 87%.46 These were higher where the study included twice daily dosing, and lower with lower frequency of dosing. It is interesting to note that the histological clearance rate was higher than the clinically observed clearance rate35,47 as reported above. In a pooled analysis of 5 studies testing higher and lower dosing regimes for BCC (not only sBCC) there was a 50% reduction in the risk of early treatment failure with the more frequent dosing regime than the less frequent.10

Side effects and disadvantages of treatment
The side effects from use of imiquimod are mainly local site reactions. In the RCTs, these tended to be complaints of erosion, ulceration and induration (in the more frequent dosing regimes) to itching, burning or pain affecting from 58% to 92% of trial participants.35,46,47,51 Prescribed or patient-initiated treatment breaks due to side effects ranged from 3% to 19%. An association was shown between severity of local site reaction and clinical response rate. The more reaction, the better the response.35,56 Despite this, few patients in any of the studies discontinued because of the inflammatory response.35,46,47,55,56

A recent study reported a case series of 24 patients (6 of whom had sBCC) examining the treatment failures and clinical characteristics associated with failure. The authors in this study emphasize the need to inform patients as to whether their prescribed use of imiquimod meets the approved uses from the drug-regulatory authority and warn of risks associated with transferring responsibility of care for a cancer to the patient rather than the physician. They advise long-term follow-up.57 A further study reports on persisting hypopigmentation following treatment.58

Cost effectiveness of treatment options
There have been few economic evaluations of the cost-effectiveness of various treatment options for non-melanoma skin cancer – even less specifically for sBCC.11 In one study in Spain the cost effectiveness of treatment with imiquimod was compared with regular excisional surgery. The findings may not be generalizable because they are based on the Spanish health care system but the principles underlying the assumptions about care such as when, where and by whom treatment is delivered provide a framework that could apply to other geographical regions. The findings indicated reduced costs per patient cured for imiquimod compared to surgical treatment in either a dermatological or non-dermatological service. Costs of primary care physicians were only involved in follow-up and any subsequent late treatment failure costs were not included.59 On the other hand, in Australia, General Practitioners (primary care physicians) tend to treat most sBCC, and with surgery,15 although there has been no economic evaluation of this care specifically for sBCC in Australia.

Other regimes and options for the use of imiquimod
More recent trials have focused on the different regimes but with only small numbers of patients, but longer follow-up. Five weeks of 5% imiquimod cream once daily with a 1-week interval was more effective but as well tolerated as the 8-week alternate week regimen for sBCC.60 Using occlusion 3 days per week appeared not to improve the success rate for sBCC (87%) over the more conventional treatment regime of...
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<th>Study</th>
<th>Design</th>
<th>Subjects</th>
<th>Intervention</th>
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<th>Adverse effects</th>
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<tr>
<td>Geisse et al(17)</td>
<td>Prospective, randomized</td>
<td>724 patients, sBCC, min 0.5 cm² area, max 2 cm diameter</td>
<td>5% imiquimod cream: 1) 5 × per wk for 6 wk</td>
<td>Clinical + histological evidence of BCC 12 wk post treatment</td>
<td>Clinical + histological clearance rate: 1) 5 × per wk: 75% (139/185) 2) 7 × per wk: 73% (130/179) 3) vehicle: 2% (6/360). Histological clearance rate: 1) 5 × per wk: 82% (152/185) 2) 7 × per wk: 79% (142/179) 3) vehicle: 3% (11/360)</td>
<td>At least one adverse event in 58% of 5 × per wk for 6 wk group: 64% of 7 × per wk for 6 wk group: 36% in vehicle group. Application site reactions most common, including itching, burning and pain</td>
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<td>3) vehicle: 6% (2/360).</td>
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<td>Schulze et al(15)</td>
<td>Prospective, randomized</td>
<td>166 patients, sBCC, min 0.5 cm² area, max 2 cm diameter</td>
<td>5% imiquimod cream daily for 6 wk</td>
<td>Clinical + histological evidence of BCC 12 wk post treatment</td>
<td>Clinical + histological clearance rate: 77% for treatment group vs 6% for vehicle group. Histological clearance rate: 80% for treatment group vs 6% for vehicle group</td>
<td>At least one adverse event in 52% of treatment group; 18% of vehicle group. Application site reactions most common, including itching and burning</td>
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<td>Beutner et al(11)</td>
<td>Prospective, randomized</td>
<td>35 patients, sBCC (28 patients, 0.5–2 cm²), nBCC (7 patients, 0.5–1.5 cm²)</td>
<td>5% imiquimod cream in one of five dosage regimens (24/35) vs vehicle (11/35)</td>
<td>Clinical and histological evidence of BCC 6 wk post treatment</td>
<td>Clearance rates: overall 83% (20/24) 1) twice daily for 10 wk: 100% (7/7) 2) once daily for 13 wk: 100% (4/4) 3) 3 × per wk: 14.5 wk: 100% (4/4) 4) 2 × per wk: 16 wk: 80% (3/5) 5) 1 × per wk: 16 wk: 50% (2/4) 6) vehicle, 16 wk: 9% (1/11)</td>
<td>Application site reactions in 92% (22/24) of treatment group, 64% (7/11) in vehicle group, including itching, erythema, papular rash and discharge. Several local reactions (erosion, induration, ulceration) observed only in twice daily and once daily groups</td>
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<td>double-blind vehicle-controlled</td>
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<td>Administration 2 wk after clinical clearance or up to 16 wk</td>
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<tr>
<td>Geisse et al(16)</td>
<td>Prospective, randomized</td>
<td>128 patients, sBCC (0.5–2 cm²), 24 patients withdrew from treatment portion of study</td>
<td>5% imiquimod cream in one of four dosage regimens for 12 wk</td>
<td>Clinical and histological evidence of BCC 6 wk post treatment</td>
<td>Clearance rates: 1) twice daily: 100% (10/10) 2) once daily: 87% (27/31) 3) 5 × per wk: 81% (21/26) 4) 3 × per wk: 52% (15/29) 5) vehicle: 19% (6/32)</td>
<td>118/128 reported at least one adverse event. Most frequently reported application site reactions included itching, pain and tenderness at application site</td>
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Table 2 Double-blind randomized controlled trials investigating 5% imiquimod cream for the treatment of basal cell carcinoma (BCC) including superficial basal cell carcinoma (sBCC)

5 times per week for 12 weeks. One study reports treatment of 96 patients with 141 lesions, 45 of which were biopsy proven sBCC mainly on the face neck or scalp. The patient group was considered to be unsuitable for or had refused surgery. They used the cream 3 days per week with multiple applications from one sachet to reduce cost. This pragmatic carefully designed study produced clinical cure rates of 83% to 87% at 6 months and 80% to 85% at 1 year despite a less frequent dosing regime.61

There is also a scientific rationale for combining surgical techniques with the use of imiquimod either to decrease the initial tumor size or to debulk it or to “mop-up” residual tumor cells following surgery on difficult body sites.62 One study reports improved results from this approach. The numbers were small (57) and included sBCC and nBCC where curettage without cautery was performed and followed one week later by imiquimod 5% cream applied 5 times per week for 6 weeks. At 1-year follow-up there were no clinical recurrences and cosmetic results were reported as excellent.63 A further prospective follow-up study also reports treating patients with biopsy confirmed BCC (unknown whether they were superficial type) using curettage followed by imiquimod. A sample of lesions were biopsied at 6 weeks after therapy, two of which showed BCC, but on average follow-up at 36 months of 101 lesions the clearance rates were 96% with favorable cosmetic outcome.64 Imiquimod has also been used as adjunctive therapy where Mohs surgery is planned for sBCC or nBCC. A report of where Mohs was undertaken for three cases of large mixed-type BCC was ceased after the removal of aggressive tumor components and the remaining sBCC received adjuvant therapy with imiquimod cream for 6 weeks showed no recurrences after a follow-up period of 20 to 34 months.65

Recurrence and long-term follow-up

At this stage, in two open label phase III studies of sBCC treatment regimes of once daily 5 or 7 times per week (n = 169) application of imiquimod 5% cream, the 2-year follow-up has been reported. The outcome measure has been a clinical evaluation rather than excision, for obvious reasons. This approach also mimics the reality of clinical practice. The treatment regime in the first study was 5 times per week application for 6 weeks (n = 182) of whom 178 completed treatment. The follow-up of this study has subsequently been published and indicates the overall efficacy of the treatment at 5 years (with all patients included) was 78% clinically cleared and 81% histological clearance. One patient who was clear at the 12-week visit died prior to entering follow-up. It was noted that if recurrences occurred in this study they mostly occurred during the first 9 months after the end of treatment. The initial response was therefore predictive of long-term outcome so these authors recommend and encourage continued monitoring of skin lesions. The second study demonstrated a clearance rate of 94% at 12 weeks post treatment and those clinically clear at 2 years were 82% with no difference between five times per week treatment and seven times per week treatment for recurrence, but the more frequent dosing increased side effects.66 A further small open study of 5-year follow-up for 55 lesions, showed of the four sBCC the long-term clearance rate was 100%.67

Other reports on imiquimod in special case sBCC

The approval by licensing authorities has focused on the use of this preparation for skin malignancies on small non-head and neck sBCC and AK. Trials are still underway for its use in nBCC, Bowen’s disease, and larger lesions, these topics are outside this review. One case report discusses success using imiquimod for a patient with multiple sBCC, after failure of treatment with 5FU and a second reports on the effectiveness of imiquimod for a single large (30 cm²) sBCC with histological assessment at 10 months post treatment.69 There are also a number of reports of case series where patients with BCNS (Gorlins syndrome) with multiple facial and trunk superficial and nodular BCC have been treated with a high level of success using imiquimod cream.70 There may also be a role for the treatment of sBCC occurring in high risk renal transplant patients with one study reporting that it appears to be safe on skin areas up to 60 cm².71

Conclusion

Long-term follow-up of blinded RCTs comparing imiquimod with other interventions are not yet available although these are ongoing. Only one RCT report including sBCC is available for 5-year follow-up and this compares cryosurgery with PDT. While awaiting final outcomes of these studies it would appear that imiquimod should not be first-line treatment for superficial BCC, as surgical excision is more effective. However, the self-treatment regimes and demonstrated efficacy so far are attractive.71 There is reasonable evidence that the use of imiquimod for small (<2 cm) superficial BCC that occur other than on the face provides outcomes only marginally less satisfactory than surgery. There would be a place for imiquimod in treating patients with frequent multiple primary lesions when access to surgery is difficult or where clinical judgment may be influenced by
patient factors as reported in some of the studies, eg, where patients may have contraindications to surgery. Cosmetic outcome may also be relevant since sBCC occurs more frequently in younger patients.

Monitoring and follow-up are important if using imiquimod. There are a substantial proportion of lesions that do not respond, and this may be because they are unrecognized as having mixed components even if biopsy is performed and patients who have one BCC are at risk of more.

Side effects that are unpleasant and unsightly such as erosion, ulceration, burning and stinging, although commonly occurring in the majority of successful treatments of sBCC, are a two-edged sword. They generally indicate a high level of immune response and are correlated with clearance of the lesion. When patients were warned to expect these side effects, as in the trials, there was a minimal drop out because of them.

Areas for further research

More research is needed focusing on the use of imiquimod for sBCC on the face, those of larger size, the relationship between cure rates and side effects, and in the role of imiquimod as adjuvant therapy to or prior to surgical procedures especially in cosmetically sensitive areas. In addition, since imiquimod is a self-treatment regime, the outcomes of treatment in patients in rural areas with poor access to specialized care compared with other approaches would be valuable.

Disclosures

The author has acted as a consultant for 3M Pharmaceuticals.

References


