Reflectance Confocal Microscopy for Noninvasive Monitoring of Therapy and Detection of Subclinical Actinic Keratoses

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Introduction

Actinic keratoses (AK) are the most common malignant skin neoplasms and have been histopathologically classified as cutaneous squamous cell carcinoma in situ [1, 2]. Sites of predilection are areas of chronic, high-dose or intermittent sun exposure, favoring the face, neck and forearms in persons with Fitzpatrick skin phototype I–III. As a ubiquitous carcinogen, UV radiation induces the initiation and promotion of progressive keratinocyte dysplasia, often affecting large areas of skin [3]. This etiopathogenetic concept has been referred to as ‘actinic field cancerization’; AK represents one stage in the continuum from subclinical keratinocyte dysplasia to invasive squamous cell carcinoma [4, 5]. The diagnosis of AK is generally based on clinical examination. However, biopsy and histological evaluation are performed in clinically indistinct cases and if invasive squamous cell carcinoma is suspected. While visual inspection may fail to detect early subclinical changes, invasive, repeated histological analysis of areas with extensive actinic damage may often not be feasible.

In the past decade, a number of novel treatment modalities have been developed and evaluated regarding their efficacy and safety. Imiquimod (IMIQ) 5% cream...
represents the first topical immunomodulator approved for the treatment of genital warts, superficial basal cell carcinoma and AK [6–8]. For the treatment of AK, complete response rates ranging from 50 to 84% have been reported [7, 9]. Recently, new insights have been gained concerning the mode of action of IMIQ 5% cream. It acts as an agonist of toll-like receptors 7 and 8 resulting in activation of NF-κB and subsequent induction of proinflammatory cytokines including among others INF-α, TNF-α, IL-2, IL-6, IL-8 and IL-12 [10]. In this induction, a central role has been ascribed to antigen-presenting dendritic cells, responding to low threshold concentrations of IMIQ [11, 12]. Following topical application of IMIQ, the maturation and migration of epidermal Langerhans cells to regional lymph nodes has been demonstrated in vivo [10]. Summarizing current observations, the mode of action is based on the activation of cytotoxic T cells with antitumoral and proapoptotic properties and induction of the innate immune response, which – in an independent pathway via the activation of the caspase cascade – results in mitochondrial-mediated apoptosis [13, 14]. A number of clinical trials and observational analyses have shown the efficacy of IMIQ treatment for AK. While it has been demonstrated that IMIQ is able to induce a prominent inflammatory response in areas of visible AK, selected subclinical lesions may show a comparable response [15]. In the context of actinic field cancerization, this has also been referred to as the visualization of ‘subclinical lesions’. Similarly, some studies reported a cosmetic improvement of areas with actinic photodamage following treatment with IMIQ, suggesting an overall regenerative effect of this immunomodulatory treatment [16, 17].

Reflectance confocal microscopy (RCM) is a noninvasive imaging technique that allows the visualization of cellular and subcellular structures of the skin in vivo with near histological resolution. In contrast to histological evaluation, which visualizes vertical sections of the tissues, RCM obtains horizontal (en face) optical sections in gray scale. A digital camera (Viva Cam, Lucid Inc., Rochester, N.Y., USA) connected to the RCM computer obtains dermoscopic images which may then be directly correlated to RCM evaluation and guide the identification of suspicious areas within the lesion [18]. A single image allows in vivo evaluation of a 500 × 500 μm area and by scanning of the microscope composite images, up to 8 × 8 mm can be obtained. RCM has been used for the evaluation of a variety of inflammatory and neoplastic skin disorders [19–24]; recently, the clinically applicability of this method for the evaluation of AK has been investigated [25–27]. It was postulated that initial changes of epidermal morphology and cellular atypia may be visualized by RCM before becoming clinically apparent. Therefore RCM may be useful for the evaluation of actinic field cancerization and the detection of subclinical AK. Moreover, the lesions may be examined in vivo and repeatedly over time, enabling a noninvasive analysis of treatment effects by RCM.

Based on these hypotheses, the primary goal of this study was to evaluate and describe the inflammatory and regenerative pharmacodynamic changes induced by IMIQ treatment over time. The secondary aim of this study was to test the possibility of detecting subclinical AK using RCM by correlating the microscopic observations to the clinically visible immunologic response to IMIQ.

Subjects and Methods

Participants

11 otherwise healthy Caucasian volunteers (Fitzpatrick skin phototype II–III), aged between 59 and 77 years, with a clinical diagnosis of AK of the face and scalp and signs of field cancerization were recruited to participate in this study. Subjects with a history of significant other skin disease were excluded from the protocol. Clinical investigations were conducted according to the principles of the Declaration of Helsinki and consent was obtained prior to enrollment. All 11 subjects completed the study and data of all subjects were included in the analysis.

Clinical Evaluation

All participants were examined for the presence of AK following the guidelines of the visual inspection and diagnosis of skin cancer [28]. 11 skin sites clinically suspicious for AK (subsequently defined as clinical AK) and 11 perilesional, clinically uninvolved skin sites (subsequently defined as subclinical AK) were selected for RCM analysis. Clinical photographs of all skin sites were obtained using a digital camera (Sony Cybershot 7.2, Sony Corp., Tokyo, Japan; Canon 350, Canon, Tokyo, Japan) under standardized room and lighting conditions. Transparency body charts were used to outline the selected skin areas at baseline to permit colocalization during follow-up examination for the duration of the study. Similarly, highlighting of new skin areas was documented on the chart by using permanent marker pens in a color-coded fashion, correlating individual colors to selected evaluation dates.

Treatment Protocol

IMIQ (MEDA Pharma, Sweden) was applied to selected study sites at recommended intervals 3 times weekly for a total of 4 weeks. Patients were requested to document their application regimen and possible treatment-related side effects using a patient diary.
Table 1. Presence of selected features in clinical and subclinical AK sites

<table>
<thead>
<tr>
<th>Feature</th>
<th>Subclinical AK</th>
<th>Clinical AK</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Main features</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Parakeratosis/hyperkeratosis</td>
<td>○</td>
<td>●</td>
</tr>
<tr>
<td>Stratum corneum disruption</td>
<td>○</td>
<td>●</td>
</tr>
<tr>
<td>Individual corneocytes</td>
<td>○</td>
<td>●</td>
</tr>
<tr>
<td>SG – pleomorphism</td>
<td>○</td>
<td>●</td>
</tr>
<tr>
<td>SG – architectural disruption</td>
<td>○</td>
<td>●</td>
</tr>
<tr>
<td>SS – pleomorphism</td>
<td>●</td>
<td>●</td>
</tr>
<tr>
<td>SS – architectural disruption</td>
<td>●</td>
<td>●</td>
</tr>
<tr>
<td><strong>Additional features</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Solar elastosis</td>
<td>●</td>
<td>●</td>
</tr>
<tr>
<td>Increased vascularity/blood vessel dilatation</td>
<td>○</td>
<td>●</td>
</tr>
</tbody>
</table>

SG = Stratum granulosum; SS = stratum spinosum. ○ = Absent; ● = present.

**In vivo RCM Evaluation**

A commercially available in vivo RCM device (Vivascope 1500, MAVIG GmbH, Munich, Lucid-Tech Inc., Henrietta, N.Y., USA) was used for imaging of all test sites. A detailed description of this technique and the device has been published [29–31]. Each skin site was systematically evaluated by RCM; one X-Y horizontal mapping (map dimensions: 4 mm × 4 mm) was performed at each epidermal layer beginning with the stratum corneum through the entire epidermis and into the upper reticular dermis. Similarly, 4–6 individual RCM images were captured in each epidermal layer, following a standardized evaluation protocol and individually subjected to descriptive analysis during the evaluation process. Based on previously defined RCM criteria, specific study sites were evaluated for the presence or absence of RCM features of AK [25–27]. The presence of main features was indicative of AK: disruption of the stratum corneum, individual corneocytes, parakeratosis/hyperkeratosis, overall architectural disruption, nuclear atypia/cellular pleomorphism at the level of the basal, spinous and granular layer. Additional features were used as follows: spongiosis, exocytosis, solar elastosis, increased vascularity/blood vessel dilatation, lymphocyte rolling, superficial inflammatory infiltrate. Based on these criteria, the presence of nuclear atypia/cellular pleomorphism was mandatory for a diagnosis of AK using RCM. The investigator (M.U.) performed the evaluation of all test sites and was not blinded to the clinical diagnosis before initiating RCM evaluation.

**Detection of Clinical/Subclinical AK.** Areas that showed main features of AK on RCM evaluation in the presence of clinical findings were subsequently defined as ‘clinical AK’; corresponding areas that showed main features of AK on RCM evaluation in the absence of clinical findings were defined as ‘subclinical AK’.

**Therapeutic Monitoring.** To detect the morphologic changes in study sites following topical applications of IMIQ 5% cream, serial RCM evaluations were performed at selected intervals, including baseline (before treatment), 2 weeks after initiation of therapy and 4 weeks after end of treatment. The presence or absence of RCM features of AK was recorded as described above (table 1); similarly features of inflammatory response, including the presence of spongiosis, inflammatory cells, intraepidermal necrosis and superficial impetiginization or crusting, were documented. Clinical images illustrating the course of IMIQ treatment are shown in figure 7.

**Results**

**Clinical AK**

**RCM Evaluation of Clinical AK before Initiation of Treatment (Baseline)**

At the level of the stratum corneum, single detached keratinocytes seen as bright polygonal structures and clinically corresponding to hyperkeratotic scaling were visualized in all evaluation sites. At the level of the stratum spinosum and granulosum, atypical keratinocytes with cellular and nuclear pleomorphism were seen. An overall disarray of epidermal architecture with loss of the normal honeycomb pattern was visualized (11/11). At the level of the superficial dermis, blood vessel dilatation (8/11) and solar elastosis (11/11) were seen (fig. 1).

**RCM Evaluation of Clinical AK after 2 Weeks of IMIQ Treatment**

Upon clinical evaluation, a notable inflammatory response was documented in 10/11 patients. One patient (patient 1) did not show any local or systemic response but continued treatment and was followed by RCM.

RCM evaluation of AK during treatment with IMIQ showed persistent superficial disruption at the level of the stratum corneum. At the level of the stratum spinosum and granulosum, persisting cellular and nuclear atypia was seen. At the level of the dermis, bright irregular bundles were visualized, representing solar elastosis. As aspects of inflammation/immunomodulation, RCM evaluation at the level of the stratum corneum revealed aggregates of small, weakly refractive cells clinically corresponding to superficial impetiginization/crusting in 7/11 patients (not shown). In these areas, optical penetration depth was limited, partially interfering with resolution of deeper layers, impeding the visualization of cellular detail and inflammatory response in 4/11 patients. RCM evaluation at the level of the epidermis and dermis revealed large, dendritic structures of bright appearance as well as small round bright cells of 10 µm in diameter scattered among keratinocytes (6/11). Dendritic cells of two morphological subtypes were visualized by RCM: one of spindle...
shape and high reflectance, resembling melanocytes as described by Pellacani et al. [21] and the other of large shape and bright appearance, most likely corresponding to epidermal (Langerhans) or dermal dendritic cells or melanophages. At the level of the dermis, the presence of small round bright cells was visualized in a perifollicular and perivascular distribution (6/11). Blood vessels of increased diameter were seen and high blood flow was noted in vivo examination (6/11) (fig. 2).

RCM Evaluation of Clinical AK 4 Weeks after the End of IMIQ Treatment

Clinical examination showed clearance of AK in 9/11 patients; patient 1 did not to show an identifiable treatment response on clinical examination and was thus classified as a nonresponder. Another patient (patient 2) showed clearance on clinical examination but persistence of AK features (atypia of the stratum granulosum and spinosum) and inflammation on RCM examination. Overall, the stratum corneum showed a more homogeneous appearance and increased cohesiveness of corneocytes comparable to findings observed in normal, unaffected skin sites [25–27]. RCM images obtained at the level of the stratum granulosum and spinosum showed a more regular appearance of keratinocytes, with polygonal cells arranged in a typical honeycomb pattern (9/11). At the level of the dermis, the persistence of bright, irregularly configurated bundles and lace-like amorphous material suggestive of solar elastosis was seen in all patients. Superficial dermal blood vessels showed a regular appearance in the majority of patients (8/11) with no residual inflammatory infiltrate present (fig. 3).

Subclinical AK

RCM Evaluation of Subclinical AK before Initiation of Treatment (Baseline)

At the level of the stratum corneum, a cohesive layer of corneocytes with homogeneous bright reflectance and
Fig. 2. RCM morphology of clinical AK 2 weeks after initiation of IMIQ treatment.

a RCM image obtained at the stratum spinosum illustrating atypical keratinocytes and the presence of large, spindle-shaped dendritic cells (dashed circle) and small, highly refractive structures (arrows). 

b RCM image obtained at the suprabasal layer showing increased brightness of suprabasal keratinocytes. The cells measure around 20 μm in diameter and have large dark nuclei and bright cytoplasm (dashed circles). Furthermore, small round bright cells of around 10 μm can be identified, which likely correspond to inflammatory cells (arrows).  

c RCM image obtained at the level of the dermo-epidermal junction showing 3 large very bright cells of dendritic appearance in perifollicular distribution (dashed circles), surrounded by several round bright cells suggestive of inflammatory cells. 

d RCM image obtained at the level of the papillary dermis with presence of prominent blood vessels (BV) that appear as bright canalicular structures and showed increased blood flow on in vivo examination. Furthermore, a perivascular and perifollicular inflammatory infiltrate consisting of small round bright cells can be seen.

Fig. 3. RCM morphology of clinical AK 4 weeks after the end of treatment.

a RCM image obtained at the level of the stratum corneum showing homogeneous appearance and coherence of corneocytes. 

b RCM image obtained at the level of the stratum spinosum with regular keratinocytes arranged in a typical honeycomb pattern and visualization of typical skin folds (SF). 

c RCM image obtained at the level of the upper reticular dermis with thickened collagen bundles and moderately refractile lace-like amorphous material illustrating remaining solar elastosis.
A regular appearance was observed in 9/11 patients. 2/11 patients showed slight superficial stratum corneum disruption with single detached corneocytes. The stratum granulosum showed polygonal, uniform cells arranged in a typical honeycomb pattern in the majority of patients (9/11). At the level of the stratum spinosum, discrete cellular and nuclear atypia of the keratinocytes was visualized, resulting in focal loss of the honeycomb pattern (dashed circle). Arrows indicate nuclear polymorphism. Furthermore, small canalicular bright structures morphologically corresponding to dilated blood vessels were seen in 3/11 patients (fig. 4).

**RCM Evaluation of Subclinical AK 2 Weeks after Initiation of IMIQ Treatment**

Clinical evaluation revealed a notable inflammatory response in 10/11 patients. Patient 1 did not show any signs of clinical response. RCM evaluation of subclinical sites showed persistence of RCM features of AK in all lesions at the level of the basal and spinous layer with atypia of keratinocytes and nuclear pleomorphism. In contrast to the baseline evaluation, superficial disruption and impetiginization/crusting was seen, partially interfering with optical penetration and impeding the visualization of inflammatory response in 2/11 lesions. Variable degrees of inflammation were detected within the epidermis and dermis, seen as small bright round cells and large dendritic cells in 8/11 patients. The presence of single denticulated large cells may likely correspond to apoptotic keratinocytes. At the level of the suprabasal layer cells two characteristic cell types were frequently observed: one larger type of approximately 15 μm in diameter and a dark nucleus surrounded by a highly refractive cytoplasm consistent with pigmented basal cells; scattered small bright cells of approximately 10 μm corresponding to inflammatory infiltrate were visualized. Similarly, numerous small bright round cells were detected around hair follicles and were
interspersed in between collagen bundles. Large canalicular spaces filled with small round cells of medium reflectance and increased flow on in vivo examination were visualized that correspond to dilated and elongated blood vessels (8/11) (fig. 5).

**RCM Evaluation of Subclinical AK 4 Weeks after the End of IMIQ Treatment**

Upon clinical evaluation, resolution of inflammation was noted in all patients with a previously documented treatment response. The nonresponder (patient 1) did not show significant changes when compared to the baseline visit in both clinical and RCM evaluation. Patient 2 showed no clinical findings of AK while RCM features of AK and inflammation persisted. The stratum corneum appeared homogeneously bright and showed cohesion of tight corneocytes in 10/11 patients. At the level of the stratum granulosum and spinosum, polygonal cells of regular appearance and homogeneous arrangement were seen in the majority of patients (8/11), resulting in a typical honeycomb pattern. Interpretation of RCM data was limited in 1 patient due to poor image quality. At the level of the dermis, persistence of bright thickened bundles was observed (fig. 6).

**Discussion**

The incidence of AK has been consistently rising, now accounting for the second most common cause for dermatology consultations in the United States [32]. AK typically arise in areas of chronic sun exposure of the face, neck and forearms and represent one stage in the continuum from subclinical keratinocyte dysplasia to invasive squamous cell carcinoma. The diagnosis of AK is generally based on clinical inspection, and studies have shown positive predictive values for clinical diagnosis between 81 and 94% when compared to histopathologic examination [33, 34]. Biopsy with histological evaluation is not routinely performed in the diagnostic work-up for...
AK, nor is histology generally performed to evaluate treatment efficacy, with the exception of selected outcome studies and for the detection of residual, recurrent or invasive disease. In an attempt to increase diagnostic accuracy with noninvasive imaging technologies, a number of adjunct diagnostic devices have been evaluated for their clinical applicability. Among others, recent studies reported high sensitivity and specificity rates for diagnosis of AK by RCM in vivo [26, 27].

Similarly, the access to noninvasive, topical treatment modalities has greatly facilitated the management of AK [35]. Among them the topical immunomodulator IMIQ...
is now widely used in dermatologic therapy. Its mode of action implies the induction of the innate immunity resulting in a selective antitumoral immune response through the activation of dendritic cells, cytotoxic T lymphocytes and proinflammatory cytokines. RCM has been used for monitoring and evaluation of treatment efficacy of basal cell carcinoma and lentigo maligna melanoma during topical IMIQ therapy [36–38]. To our knowledge, no systematic RCM evaluation of treatment response of AK to IMIQ has been performed.

The primary objective of this study was to evaluate and describe the dynamic morphologic changes induced by IMIQ treatment over time. Furthermore we aimed to test the possibility of detecting subclinical AK using RCM in order to confirm this hypothesis by the clinically visible response to IMIQ.

Following baseline evaluations of clinical and subclinical AK, all patients participating in this study were asked to apply IMIQ 5% cream in a defined study area 3 times weekly for a total of 4 weeks, following established dosing schemes. Neoplastic, inflammatory, regenerative and vascular changes were documented in both clinical and subclinical AK sites using RCM evaluation parameters for AK as published previously [26, 27].

RCM was able to identify subclinical AK by visualization of cellular and nuclear atypia within the spinous cell layer. After 2 weeks of treatment, RCM morphologic features of inflammatory response were detected by RCM in vivo and included the presence of small bright cells and two types of dendritic cells scattered among keratinocytes and the superficial dermis. The spindle-shaped, bright dendritic cells most likely correspond to melanocytes [39], whereas the plumb bright dendritic cells may correspond to Langerhans or dermal dendritic cells, which have previously been shown histologically to belong to the inflammatory infiltrate in IMIQ-treated lesions [40]. The presence of the melanocytes may be explained by the chronically sun-damaged skin with mottled hyperpigmentation that is often associated with AK. Furthermore, the detection of denticulated large cells within the epidermis may correspond to apoptotic keratinocytes, an observation that has previously been described in the mouse model [10].

Furthermore, RCM was able to detect residual atypia in 2 patients after end of treatment without any clinical aspects of AK. These findings were suggestive of incomplete clearance of AK, subsequently prompting a second cycle of IMIQ therapy. In that context, it may be noted that following noninvasive therapy, a histopathologic control of clearance is not routinely performed and not feasible. By using RCM for evaluation of treatment efficacy, the early, noninvasive detection of residual atypia may ultimately decrease recurrence rates.

Limitations of the technique include the time needed for image acquisition in large anatomic areas as well as inherent optical limitations interfering with resolution of cellular details. The optical resolution and penetration is especially limited in hyperkeratotic lesions such as squamous cell carcinoma, verrucous lesions or hyperkeratotic AK and may interfere with image quality and correct diagnosis may often not be obtained by RCM. While the technique is easily learned, the results remain strongly user-dependent, and reproducibility of results may vary between investigators.

In conclusion, our findings indicate that the use of RCM allows the noninvasive visualization and monitoring of sequential pharmacodynamic changes during IMIQ therapy over time. Thereby, RCM may expand our insights into the mode of action of this immunomodulatory treatment and facilitate the evaluation of treatment efficacy. Furthermore, RCM may aid in the detection of subclinical AK in the setting of field cancerization, potentially increasing diagnostic accuracy compared to clinical evaluation alone. The adjunct use of noninvasive optical techniques may allow the early identification of affected skin sites, substantiate the incentive for therapy and ultimately increase therapeutic efficacy.

Conflict of Interest

M. Ulrich was awarded a research grant by MEDA Pharma GmbH, Sweden; M. Ulrich, S. Astner and E. Stockfleth are lecturers for MAVIG GmbH, Germany; E. Stockfleth is a consultant for MEDA Pharma GmbH.

References


