Successful Treatment of Genital Warts with Ingenol Mebutate Monitored with Optical Coherence Tomography and Reflectance Confocal Microscopy

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Ingenol mebutate (IM) is approved for the treatment of actinic keratosis and induces cell death in precancerous lesions. The efficacy of IM in the treatment of genital warts was investigated in a therapy-refractory patient. The 74-year-old male was treated with IM gel for three consecutive days. Treatment course and efficacy were evaluated by clinical inspection and non-invasive diagnostics namely optical coherence tomography (OCT) and reflectance confocal microscopy (RCM). Within 24 to 48 hours IM induced a strong local inflammatory reaction. One week later a complete response was observed. OCT and RCM showed a strong reaction after treatment with erosions, swelling of cells, and a subepidermal dark band in representative lesions. IM has the advantage of a short treatment period in contrast to other topical treatments and shows a promising clinical outcome. Larger studies are needed to validate the data. (Ann Dermatol 31(4) 434 ~ 437, 2019)

-Keywords-
Condylomata acuminata, Microscopy, confocal, Neoplasms, Optical coherence tomography

INTRODUCTION
Infections with human papillomavirus (HPV) are a modern pandemic with millions of people being infected each year. The 170 different types of this double-stranded DNA virus are divided into low- and high-risk virus classes according to their oncogenic potential. The high-risk subtypes are commonly found in cervical, penile, or anal dysplasias with the potential to progress to carcinomas after a long-standing infection. Low-risk virus subtypes can lead to the development of genital warts, which are biologically benign but very bothersome and stigmatizing for the patient. Moreover HPV is highly contagious; therefore genital warts are a very common manifestation of HPV infection in sexually active adults. Sexual contact with an HPV-infected individual results in a 75 percent chance of contracting the virus, with additional risk factors being immunosuppression, unprotected intercourse, smoking, or a history of sexually transmitted diseases.

For the treatment of genital warts various topical medications including 5-fluorouracil, sinecatechins, trichloroacetic acid, and imiquimod are available. Other treatment options include cryosurgery, laser, electrocautery, or surgical excision, but with higher risk of scarring. Recently, ingenol mebutate (IM) has been approved as a new agent for topical treatment of actinic keratosis. IM is one of the active compounds of the plant *Euphorbia peplus* L. The
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hydrophobic diterpene ester called ingenol-3-angelate was identified as an active ingredient. Usage of wild E. peplus for treatment of various medical conditions is known from ancient times.

The objective of our investigation was the initiation of an immune response to the genital warts through topical application of IM. We hypothesized that IM would trigger apoptosis and necrosis of the affected cells and additionally induce an antigenic immune response.

Modern imaging techniques such as optical coherence tomography (OCT) and reflectance confocal microscopy (RCM) have proven useful in practice for the noninvasive diagnosis of non-melanoma skin cancer and treatment monitoring.

Reaction and response to treatment were mainly evaluated by clinical examination, while OCT and RCM were used to monitor the response, similar to monitoring the therapeutic success of IM gel in actinic keratosis.

**CASE REPORT**

In this case report, we evaluated a 74-year-old male patient who had several unsuccessful standard treatments previously. The patient presented with 15 genital warts in the area of the radix penis and scrotum (Fig. 1). The patient was treated at the Department of Dermatology, University Hospital of Munich (Ludwig Maximilian University, LMU), after obtaining written informed consent for the individual off-label use of topical ingenol mebutate in genital warts. A single dose of 0.47 g IM gel (150 μg/g, daily dose: 70.5 μg IM; Picato®; LEO Pharma A/S, Ballerup, Danmark) was applied topically for three consecutive days in a predefined treatment area of 25 cm².

The site of infection was evaluated and monitored clinically as well as documented via OCT and RCM on baseline and day 4, 7, and 21 after initial treatment. The initial clinical evaluation revealed 15 condylomata acuminata with typical features of papillomatous or verrucous tumors or plaques with brownish-red coloration.

**Non-invasive imaging**

For non-invasive imaging the following commercially available devices were used: a Fourier-domain OCT system (Vivosight®; Michelson Diagnostics, Kent, UK) with a 1,310 nm multi-beam laser providing a penetration depth of 1.5 to 2 mm and a resolution of 5 to 7.5 μm and an in-vivo RCM device (Vivascope® 1500; Lucid-Tech Inc., Henrietta, NY, USA; Mavig GmbH, Munich, Germany) with a penetration depth of about 250 μm and a resolution of 1.25 to 2.5 μm.

OCT and RCM imaging was performed in one defined target lesion before and after seven and 21 days after treatment with IM gel.

**Clinical evaluation**

IM induced an inflammatory reaction after 24 hours in the treated area. This reaction lasted up to seven days with decreasing intensity (Fig. 1).

The observed local reactions were temporary and similar to previously described reactions seen after application in actinic keratosis.

Erythema, flaking/scaling, swelling and erosion were observed between day two and four. Additionally crusting, vesiculation and pustulation were seen.

In the patient the warts disappeared completely and were not detectable even after three weeks, neither by clinical inspection (Fig. 1) nor by non-invasive imaging (OCT or RCM) (Fig. 2).

Three weeks after application of IM gel, the skin normalized without any sign of HPV infection (Fig. 1). A residual erythema was found in the patient after 21 days (Fig. 1D). No systemic side effects were observed. Post-treatment erythema was reported by the patient, but no other adverse event.

Six months after the treatment the patient was still in remission, so it can be assumed the treatment was successful.

**OCT and RCM**

Prior to treatment, the genital warts presented as papillomatous tumors with a broadening of the epidermis and...
sometimes keratotic changes including signal-intense dots intratumorally in OCT (Fig. 2).

In RCM the genital warts showed aggregated hyperplastic nodules in the Vivablock® (Mavig GmbH) mosaic overview (5×5 mm) of superficial layers and in the single RCM image (0.5×0.5 mm) a broadened epidermis with signal-intense bright epidermal cells and hypervascularization (Fig. 2).

Four days after treatment start, OCT showed flattening of the epidermis and disappearance of layering. A signal-poor dark band sublesionally was characteristic and consistent with inflammatory reactions and edema (Fig. 2).

On day four multiple bright cells spreading throughout the epidermis were present in RCM imaging compatible with inflammatory cells, which showed disruption of cells and prominent vessels within the papillae (Fig. 2).

Three weeks after treatment, OCT imaging showed a still prevailing subepidermal signal-poor dark band. For the publishing of the photographic materials a patient’s consent form has been received.

**DISCUSSION**

In the conventional topical therapies of genital warts the treatment success depends mainly on the adherence of the patients. Due to the often long-term treatment course the patients commonly fail to maintain proper procedure or develop uncomfortable side effects like pain or burning sensations.

The common treatment options for genital warts include the following therapies: Podophyllotoxin shows high recurrence rates after clearance13 while imiquimod has lower recurrence rates14. The application of sinecatechins ointment and trichloroacetic acid has led to moderate clearance rates15.

Cryosurgery, electrosurgery, and scissors excision are additional invasive treatment options16,17. Particularly the topical drug treatments are associated with inconveniences for the patients as long-lasting, high-frequent application with partially severe side effects. Therefore the treatment with IM could serve as an alternative for the therapy of genital warts.

In contrast to the recently published case report of Braun et al.18, which described the single spot-like use of the 500 μg/g formulation on multiple condylomata acuminata, the repetitive application of the 150 μg/g formulation leads to less severe local reactions including burning sensation and pain.

The mechanism of action of IM is not yet fully understood, but latest studies suggest a dual mode of action via both cellular necrosis and neutrophil-mediated antibody-dependent cellular cytotoxicity19. It was shown that within one to two hours of application, the exposure of cancerous cells to IM induces cellular necrosis20. IM gel destroys epidermal cells within hours and induces the production of antibodies that result in neutrophils targeted to kill any residual dysplastic epidermal cells, which is important for preventing relapses following treatment19. Therefore, in contrast to more invasive approaches the herein mentioned therapy has potentially less side effects like scarring. In comparison to hitherto available topical remedies the treatment with IM has the advantage of a reduced treatment time.
In summary, IM may represent a novel treatment option for refractory genital warts. Even in a pretreated patient it showed a good clinical response and lasting results. Larger prospective randomized controlled trials are needed to objectify and evaluate this clinical concept.

**CONFLICTS OF INTEREST**

MR has received speaker’s and advisor’s fees by Galderma and MEDA Pharma GmbH & Co. Kommanditgesellschaft, BMCE none, MVH none, YH none, TR has received speaker’s and advisor’s fees by Galderma; CB has received speaker’s and advisor’s fees by Almirall-Hermal, Biofrontera, Galderma, and LEO Pharma A/S. TVB and CB have received a research grant by LEO Pharma A/S. TVB has received speaker’s fees by Almirall-Hermal. The OCT Vivosight® device used in this study has been provided by Michelson Diagnostics, UK, and the RCM Vivascope 1500® device used in this study has been provided Mavig GmbH, Munich, Germany.

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