**Squamous Cell Carcinoma as an Important Differential Diagnosis of Erosive Pustular Dermatosis of the Scalp**

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Abstract

Erosive Pustular Dermatosis of the Scalp (EPDS) is a rare inflammatory disorder with unknown etiology characterized by pustules, erosions, and crusted lesions on the scalp. The diagnosis is based on the clinical and histological presentation. In addition, Squamous Cell Carcinoma (SCC) should be considered as an important differential diagnosis. A 78-year-old man presented with crusted, non-healing, pruritic scalp lesions and erosions that had been occurring for 6 months. He was referred to us with suspected SCC, which could be excluded by multiple biopsies.

The clinical and histopathological features led us to the diagnosis of erosive pustular dermatosis of the scalp. The patient was treated with 10% salicylic ointment, topical and systemic antibiotics and topical steroids. The clinical condition promptly clears under this therapy.

**Keywords:** Erosive pustular dermatosis of the scalp; squamous cell carcinoma;

Abbreviation

Erosive pustular dermatosis of the scalp: EPDS

Squamous cell carcinoma: SCC

Introduction

Erosive Pustular Dermatosis of the Scalp (EPDS) is a rare skin disease with nonspecific clinical and histopathological findings. The pathogenesis is not well known, but some predisposing factors such as local trauma, topical retinoid acid or fluorouracil, varicella zoster and prolonged exposure of a bald scalp to UV light have been reported [1,2]. The clinical findings usually consist of pustules, erosions, and crusted lesions on the scalp. An appropriate clinical presentation and compatible histological pattern are required to diagnose the patient with EPDS. Possible clinical differential diagnoses are Squamous Cell Carcinoma (SCC), perifolliculitis abscedens et suffodiens, folliculitis decalvans, dermatitis artefacta, pyogenic granuloma, IgA pemphigus, tinea, pustular psoriasis and cicatricial pemphigoid [1-3]. Although the histological pattern is non-specific, biopsies are required to exclude severe differential diagnoses, especially SCC. This rare skin disease is more common in the elder population and affects more females than males [2,3]. It is difficult to treat and tends to relapse.

Case reports

A 78-year-old man presented pruritic crusted lesions on the scalp for 6 months. Treatment with different topical cortisone preparations was unsatisfactory. He had a previous history of hypertension, syringomyelia and spinal stenosis. Mild leukocytosis (10.2/nl) and elevated C-reactive protein (23 mg/l) were observed. Tests for electrolytes as well renal and hepatic parameters were normal. Physical examination showed several crusted lesions and erosions on the scalp (Figure 1).

At first, squamous cell carcinoma was suspected and skin biopsies were taken. This differential diagnosis was excluded by histological examination.

By microbiological examination Staphylococci (coagulase negative) were found, whereas fungal cultures were negative.

Histology from the scalp showed ulcerations with scale-crusting and superficial edema. In the dermis there were diffuse mixed cell infiltrates composed of lymphocytes, granulocytes and plasma cells. In the deep dermis scarring was seen (Figure 2). Another biopsy revealed narrowed epidermis overlying moderate mixed cell infiltrates in the dermis with exocytosis of inflammatory cells and scarring (Figure 3). Periodic Acid-Schiff
(PAS) stain was negative for mycological organisms. Direct immunofluorescence was negative for IgG, IgM, IgA and C3 depositions.

The treatment was started with 10% salicylic acid and Aureomycin (chlorotetracycline) ointment, and then continued with topical cortisone (Advantan) in association with antiseptic spray (Octenisept). Erosions were treated by antiseptic gel (polyhexanide gel) and Adaptic (wound dressing). In addition Augmentin 875/125 mg (amoxicillin and clavulante) was given as oral therapy twice daily. This resulted in improvement after 2 weeks.

The clinical presentation of EPDS consists of erosive, pustular, and crusted lesions resulting in scarring alopecia. The number of pustules can vary remarkably, and in some cases they are not present [5,3].

Upon histological examination, parakeratosis, hyperkeratosis, epidermal atrophy, inflammatory infiltrates, and occasional subcorneal pustules may be found [6,7]. Although the histopathology is non-specific, biopsies are obligatory to exclude severe differential diagnoses [5].

The clinical differential diagnosis includes squamous cell carcinoma, perifolliculitis abscedens et suffodiens, folliculitis decalvans, dermatitis artefacta, pyogenic granuloma, IgA pemphigus, Tinea, pustular psoriasis and cicatricial pemphigoid [7-9].

A number of patients with EPDS have previous squamous cell carcinoma of the skin. This severe differential diagnosis of EPDS has to be excluded carefully by skin biopsies [10].

The treatment of EPDS is difficult. Various treatments such as topical corticosteroids, topical and systemic antibiotics, topical calcipotriol or tacrolimus, oral isotretinoin, dapsone and zinc with various responses have been reported. Application of topical corticosteroids was shown to be effective, though relapses after discontinuation may occur [5,7,11,12].

After exclusion of other differential diagnoses, the clinical and histopathological findings led us to the diagnosis of erosive pustular dermatosis of the scalp in our patient. The treatment was started with 10% salicylic acid for cleaning the crusted areas. After that continued with topical and systemic antibiotics in association with antiseptic spray as well as topical corticosteroid. A complete resolution of the condition under therapy confirmed our diagnosis.

The result of this study showed that EPDS can be misdiagnosed with squamous cell carcinoma, which may result in treatment and unintended adverse events.

**References**

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