Marked Reduction of the Number and Individual Volume of Sebaceous Glands in Psoriatic Lesions

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Key Words
Psoriasis · Alopecia · Psoriatic alopecia · Sebaceous gland

Abstract

Background: Psoriasis is a chronic inflammatory skin disease characterized by plaques with inflammation, infiltration, hyper-/parakeratosis and desquamation. Microscopic findings in previous studies have revealed some degree of atrophy of the sebaceous glands in patients with psoriasis vulgaris and psoriatic alopecia. Objective: The aim of this study was to investigate possible changes of the sebaceous glands in patients with psoriatic plaques and especially psoriatic alopecia. Methods: Histological and stereological analyses were performed in skin specimens from involved and healthy-looking skin of 14 patients with psoriasis. Stereology detects and quantifies 3-dimensional structures ex vivo. Furthermore, the differentiation process of sebocytes of another 14 psoriatic patients was examined by immunohistochemistry of involved and uninvolved skin specimens. Results: A significant reduction of the number of sebaceous glands as well as of the volume of individual sebaceous glands was assessed in the lesional compared to the nonlesional psoriatic skin. Moreover, it was likely that sebocytes in psoriatic lesions may not differentiate properly. Conclusion: These findings indicate that the sebaceous gland may be a player and not an innocent bystander in the development of psoriatic lesions and especially of psoriatic alopecia.
The major question raised through the years was whether there is a difference in the clinical and histopathological findings between those psoriatic patients who present alopecia and those who do not develop alopecia. Interestingly, the latest studies have shown that a large majority of patients with psoriatic alopecia have atrophy or complete absence of sebaceous glands at the time of presentation [1, 5].

The sebaceous gland and hair follicle constitute the pilosebaceous unit of the skin. During late embryogenesis, developing hair follicles display several bulges (hair peg stage), one of which gives rise to the sebaceous gland and is located just above the hair follicle stem cell bulge and below the infundibulum of the developing follicle [6]. The sebaceous gland arises as an outgrowth of the outer root sheath of the hair follicle, while undifferentiated sebocytes emerge from peripheral basal cells and then move centrally as (first partially and later fully) differentiated sebocytes. Differentiated sebocytes produce and secrete lipid-rich sebum into the hair canal that empties to the skin surface [6–10].

Sebaceous differentiation is accompanied by differentiation stage-characteristic protein patterns. Figure 1 summarizes the main differentiation markers of sebocytes and their stages of expression. Finally, sebocytes have been suggested to become, under certain circumstances, subject to a kind of ‘keratinization’, i.e. express markers of keratinocytes and lose their ability to produce sebum [11–13]. This theory may effectively explain the observed changes of the sebaceous glands in psoriasis and is supported by the fact that sebocytes and keratinocytes derive from the same progenitor cells [14].

**Materials and Methods**

The study plan and the demographic and clinical data of the patients are summarized in figure 2 and table 1. The biopsy localizations are summarized in table 2.

A qualitative histological study of these biopsies was conducted to describe the presence or absence of the sebaceous glands and their number, distribution, and morphology (shape and size) in lesional and nonlesional skin. The volume of the sebaceous glands was estimated by the Cavalieri principle.

For the immunohistological study, biomarkers which recognize different differentiation stages of human sebocytes were applied. Table 3 summarizes the main antibodies used for immunohistochemical analysis.

For further details, see the supplementary materials (for all online suppl. material, see www.karger.com/doi/10.1159/000445942) (fig. 2; tables 1–3).
Results

Histology

Conducting estimates of the volume of the sebaceous glands, 3 of the 14 patients were found to have no visible gland tissue in the lesional skin, although there were definitely sebaceous glands in biopsies of their nonlesional skin. From the biopsies of the 11 patients who had visible gland tissue in the lesional skin, 5 originated from the back and 6 from the scalp. Figure 3 demonstrates the obvious absence or shrinkage of sebaceous glands in the lesional skin.

Table 1. Characteristics of the study population (n = 14)

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>Mean ± SD</th>
<th>Median (IQR)</th>
<th>n (%)</th>
</tr>
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<tbody>
<tr>
<td>Age, years</td>
<td>53.1 ± 12.8</td>
<td>53.5 (45–61)</td>
<td>7 (50.0)</td>
</tr>
<tr>
<td>Male gender</td>
<td></td>
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<tr>
<td>Sebaceous gland volume (psoriatic skin), mm³</td>
<td>0.018±0.030</td>
<td>0.004 (0.001–0.024)</td>
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</tr>
<tr>
<td>Sebaceous gland volume (healthy skin), mm³</td>
<td>0.057±0.053</td>
<td>0.035 (0.027–0.073)</td>
<td></td>
</tr>
<tr>
<td>Thickness of psoriatic epidermis, mm</td>
<td>332.7±92.8</td>
<td>332.0 (260.0–400.7)</td>
<td></td>
</tr>
<tr>
<td>Thickness of healthy epidermis, mm</td>
<td>80.5±22.3</td>
<td>80.9 (61.4–94.0)</td>
<td></td>
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</tbody>
</table>
Stereological Volume Estimation

The volume of the sebaceous gland tissue is shown in figure 4, and the mean volume in lesional skin was on average approximately 20% of that of healthy controls (0.018 vs. 0.057 mm³; p = 0.001). When only considering the biopsies in which both lesional and nonlesional skin contained sebaceous gland tissue (fig. 5), the volume of sebaceous gland tissue was approximately 25% of that of healthy controls (0.023 vs. 0.062 mm³; p = 0.003). Figure 6 shows schematically, in a point diagram, the differences in sebaceous gland volume between lesional and non-lesional skin. The differences in and the statistical significance of sebaceous gland volume between male and female patients between the back and the head are summarized in online supplementary table 1.

Immunohistochemistry

Out of the 14 patients, 1 showed a clear difference between lesional and nonlesional skin with a stronger expression of the early sebaceous differentiation marker cytokeratin 7 (CK7), the advanced sebaceous differentiation marker epithelial membrane antigen (EMA), and the late differentiation marker epithelial sialomucin (ESM) in the lesional skin (fig. 7). Two patients showed a slight difference with CK7, EMA, and ESM being again more strongly expressed in the lesional compared to the nonlesional skin. The rest of the patients (11/14) showed no obvious difference between lesional and nonlesional skin (online suppl. table 3). The results suggest a possible tendency of sebocytes to be less differentiated in the lesional skin.

One of the 14 patients, the same patient mentioned above, showed a stronger expression of the proliferation marker Ki67 in the lesional skin than in the nonlesional skin (fig. 8). The rest of the patients (13/14) showed no obvious difference between lesional and nonlesional skin (online suppl. table 3). The results are controversial, since a higher rate of proliferation in psoriatic skin is widely known.

No differences were found between lesional and nonlesional skin regarding the expression of the marker of keratinocyte differentiation involucrin (fig. 8; online suppl. table 3).
**Fig. 3.** Examples of the absence and shrinkage of sebaceous gland tissue in the lesional compared to the nonlesional psoriatic skin.
The results of this study clearly demonstrate a reduction of sebaceous gland tissue volume in the lesional skin of psoriasis patients, confirming previous studies [1–4]. This finding is compatible with common clinical experience, since it is well known that the psoriatic plaque is characterized by thickness, dryness, and scaling, which are major clinical findings of sebostasis. The question which arises, namely ‘does this finding indicate a role of sebaceous gland changes in the pathogenesis of psoriasis?’, belongs to the hen and the egg principle, i.e. do pathological sebaceous glands cause the reactive thickness and dryness of the epidermis or is this reduction of sebaceous glands an epiphenomenon of the epithelial pathology?

In this study, four major differentiation markers of sebocytes were used. These markers have been mentioned in previous studies as possible or definite differentiation markers of sebocytes in vitro and ex vivo. However, this study definitely adds to these contributions by clearly showing the function of these four differentiation markers, which provides a useful tool in the study of sebaceous differentiation in the future. It should be mentioned that CK7 is presented as a definite early differentiation marker and ESM as a definite late differentiation marker [7, 15–17]. Regarding EMA, several opinions suggest that this antigen is expressed by all sebocytes during the entire differentiation process, so it cannot be exactly distinguished from the other two differentiation markers [7, 16]. This means, practically, that sebocytes in all three differentiation stages may express EMA, including the ones that express CK7 or ESM. Thus, while this study cannot definitely suggest that EMA is expressed in a certain differentiation stadium only, it suggests that it is expressed either by all sebocytes or by the advanced differentiated sebocytes. Furthermore, a discrimination of the continuous differentiation process of sebocytes between three stages was made for the needs of this study; some previous studies have made up to five-stage discrimination [7]. Ki67 did not reveal any problems; it is a known proliferation marker used to visualize the proliferation of sebocytes in the basal layer. Sebocytes can proliferate and differentiate concomitantly; only undifferentiated sebocytes mostly proliferate, whereas mature ones only differentiate [18].

**Discussion**

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A further hypothesis was that the abnormal differentiation process of sebocytes in psoriatic lesions may represent a squamous keratinization able to induce inflammation, as shown in experimental settings [13, 19]. It is well established that sebocytes are highly specialized, sebum-producing epithelial cells that release their content by rupture of the cell membrane and cellular degradation with the function of holocrine secretion. Sebocytes are most commonly found in the skin in association with hair follicles (forming the pilosebaceous unit) where they arise from undifferentiated hair follicle epithelial cells. Sebum forms an integral component of the epidermal

**Fig. 7.** CK7 as an early differentiation marker was more strongly expressed in the lesional skin of patient 3. EMA and ESM as middle and late differentiation markers were more strongly expressed in the nonlesional skin of the same patient. This patient shows a lack of mature sebaceous differentiation in the lesional psoriatic skin, a finding that could explain the reduced sebum production at the psoriatic lesion.
Fig. 8. Slightly higher expression of Ki67 in the lesional skin compared to the nonlesional psoriatic skin. There were unclear differences between the expression levels of involucrin (INV). This picture shows a possible elevated rate of proliferation in the lesional skin.
On the other hand, Kure et al. however, this is a nontransient scarring disease. Brocq; however, this is a nontransient scarring disease. Braun-Falco et al.

These cells produce a proliferative population of sebocytes that, in turn, differentiates to produce the lipid/sebum-producing cells [20, 21]. The derivation of sebocytes from hair follicle keratinocytes suggests the possibility that there can exist a signal that forces the hair follicle keratinocytes to turn the normal proliferation/differentiation process toward keratinocytes instead of sebocytes and produce more keratinocytes instead of providing sebocytes to the sebaceous gland [13, 19]. This theory does not explain, however, where this signal of general squamous differentiation of the dermis and epidermis comes from. Furthermore, it does not provide any information about the sequence of events or whether the squamous differentiation of sebocytes comes before the acanthosis and hyperkeratosis of the epidermis or whether the pathology of the epidermis comes first and then, as a consequence, the atrophy of the sebaceous glands, or a signal, forces all parts of the skin to a squamous differentiation.

A recent study by Kamp et al. [22] on patients with hidradenitis suppurativa/acne inversa revealed a very similar result: significantly more atrophy of the sebaceous glands in the involved skin compared to that of healthy controls. This was the first stereological volume estimation of sebaceous glands in a skin disease. Braun-Falco et al. [23] have also shown an absence or decrease of sebaceous glands in a histological study of pseudopelade Brocq; however, this is a nontransient scarring disease. On the other hand, Kure et al. [24] found in a later stereological study that lobulation of the sebaceous glands was increased in androgenetic alopecia with a consequent noticeable increase of the sebaceous gland area, while the lobule volume remained unchanged. These findings are very interesting in combination with the findings of the present study in psoriasis because they raise the question of what these diseases have in common, since they both result in an excessive change of sebaceous gland volume, although the sebaceous gland is considered not to be the primary target of these disorders. At least psoriasis and hidradenitis suppurativa/acne inversa seem to have some characteristics in common, such as local hyperproliferation of keratinocytes, genetic background, chronic inflammation, and, as a consequence, good response to antitumor necrosis factor agents and other biologics and increased risk of metabolic disease as comorbidity. In addition, both smoking and increased body mass index impair the clinical picture of both diseases. However, several dermatological diseases share all or some of these characteristics. In fact, psoriasis and hidradenitis suppurativa/acne inversa are clinically very different dermatological diseases that do not even belong in the same classification subgroup. On the other hand, one could raise the question of whether it is simply that sebaceous gland atrophy occurs in all dermatological pathologies, such as when the epithelium is ‘ill’ and inflamed it includes the sebaceous gland. A good and simple way to confirm this is to estimate stereologically the sebaceous gland volume in different skin diseases, not only in dermatoses and chronic inflammatory diseases but also in other skin pathologies, such as epithelial skin cancer (basal cell carcinoma or squamous cell carcinoma). It would be interesting to add further comparisons with completely healthy skin in order to investigate whether the sebaceous glands are smaller in psoriasis. Such a study would entail the ethical issue of using skin biopsies from a healthy population.

The findings of this study could be a stimulus for re-examining the role of sebaceous glands in psoriasis. For example, the existing theory of psoriatic alopecia being caused by the psoriatic plaque (hyperkeratotic plaque) strangling the hair follicle can be revised for the first time. Our results suggest that psoriatic alopecia is probably developed due to a secondary abnormality of sebaceous glands rather than to the mechanical cause that has hitherto been proposed. On the other hand, however, the existence of psoriasis palmoplantaris, which selectively affects the only areas of the human body lacking hair follicles and sebaceous glands, may indicate that the involvement of sebaceous glands in psoriasis is a secondary phenomenon [25, 26].

In this project, the application of stereological principles in psoriasis revealed a clear reduction of sebaceous gland volume in the diseased skin. Further studies should investigate and expand these results to other skin diseases to determine whether our findings are isolated or a result of a common, more general pathology. Our findings suggest a new approach in the effort to elucidate the complicated cascade in the pathogenesis of psoriasis, and they implicate – for the first time – not only the epidermal/dermal layers but also the appendages of this well-known yet not fully understood disease.
Statement of Ethics

The study was approved by the Ethics Committee of the Charité – Universitätsmedizin Berlin, Freie Universität Berlin.

Disclosure Statement

The authors have no conflicts of interest to declare.

References


