Retinoids Strongly and Selectively Correlate With Keratin 13 and Not Keratin 19 Expression in Cutaneous Warts of Renal Transplant Recipients

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Objective: To compare the expression of keratin (K) 13 and K19 in cutaneous warts of renal transplant recipients (RTRs) and immunocompetent individuals (ICIs).

Design: Retrospective, nonrandomized immunohistochemical study.

Patients and Methods: Specimens from cutaneous warts of RTRs and ICIs were retrieved from the archives of the Department of Pathology, University Medical Center St Radboud, Nijmegen, the Netherlands. Twenty-one warts from RTRs and 21 from ICIs were examined. Nine RTRs (10 specimens) received either systemic acitretin or topical all-trans retinoic acid, and their effect on both keratins was assessed.

Main Outcome Measures: Frequency and expression patterns of K13 and K19 in warts of RTRs vs ICIs and the effect of retinoids.

Results: A significantly higher percentage of warts of RTRs expressed K13 compared with warts of ICIs (86% vs 14%, 18 vs 3 cases, respectively; P < .001). In warts of RTRs, retinoid treatment correlated significantly with a particularly strong, segmental K13 expression pattern, which we termed zebroid. Without use of retinoids, K13 was mostly restricted to suprabasal single cells. Keratin 19 was absent in all warts of both patient groups.

Conclusions: Retinoids strongly correlate with K13 in a characteristic zebroid pattern in warts of RTRs, making K13 a sensitive marker for retinoid bioactivity in skin (lesions) of RTRs. In non–retinoid-treated RTRs, K13 is also frequently found in warts but without the dramatic zebroid pattern noted in retinoid-treated warts.

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Epithelial keratins comprise a heterogenous group of acidic (type I) and neutral-to-basic (type II) proteins. As a general rule, they are coexpressed in specific pairings, each pair consisting of a type I and a type II keratin (K). For instance, in normal adult skin, keratin pairs K5/K14 and K1/K10 predominate in the basal layer and the suprabasal compartment, respectively.1

The type I keratins K13 and K19 are usually expressed separately in adult epithelia. Combined expression occurs only in fetal skin. Both keratins are thought to be absent in normal skin of adults except at certain body sites, such as the penile foreskin, which still contains K13. Furthermore, K13 is abundantly present in adults in internal stratified epithelia and is associated (with terminal differentiation) with suprabasal expression.2-4 Expression of K19 in adults is found in simple epithelia, such as most glandular epithelia.

In the past few years, several murine and in vitro studies have demonstrated that vitamin A and its derivates (retinoids) are important regulators of epidermal differentiation and affect keratin gene expression. In cultured keratinocytes, the induction of an embryonic type of differentiation by retinoids with reinduction of K13 and K19 expression has been well documented.5,6 In vivo topical application of retinoids on photo-aged human skin also showed induction of K13.7

Although retinoids are used as chemopreventive agents for inhibiting skin cancer in renal transplant recipients (RTRs),8,9 to the best of our knowledge there are no studies regarding the effects of retinoids on keratin expression in the skin or skin lesions of these patients.

Renal transplant recipients develop multiple warts and skin neoplasms. Immunosuppressive treatment, sun exposure, and viral infection with human papillomavirus are all implicated in the etiology of cutane-
The present study shows that RTR-associated warts in contrast to warts in normal ICIs indeed show pronounced K13 expression. This suggests that altered keratin expression may reflect an important molecular event inherent in the malignant degeneration of warts in RTRs. Furthermore, this K13 expression in warts of RTRs strongly correlates with retinoid therapy, but, in contrast to findings in animal studies and in cultured human keratinocytes, we could not demonstrate an effect of retinoids on K19 expression in these patients. Retinoid-related K13 expression in epithelial skin lesions of RTRs displays a highly characteristic pattern, which we termed zebroid, making K13 a useful marker for evaluating the effect of retinoid treatment in these patients.
esophagus was used as a positive control and showed strong diffuse suprabasal staining.

Keratin 19 immunostaining was also localized in the cytoplasm. Eccrine ducts and sweat glands, serving as internal controls, showed marked positivity.

**EXPRESSION OF K13 AND K19 IN WARTS OF RTRs VS ICIs AND EFFECTS OF RETINOID TREATMENT**

There was a significant difference in K13 expression between warts of RTRs and warts of ICIs (P < .001). A high percentage of warts of RTRs (86%, 18 cases) showed K13 expression, whereas in benign warts of ICIs almost all lesions were negative except for 3 (14%) of 21 with suprabasal single-cell positivity (Table 1 and Figure 1). This statistical difference in K13 positivity remained when we excluded retinoid-treated patients: 82% K13 positivity in non–retinoid-treated RTRs vs 14% in controls (P < .001). Besides number of positive specimens, the proportion of positive lesional cells also differed and was more pronounced in warts of RTRs: the 3 positive warts of ICIs showed only suprabasal single-cell positivity (Figure 2); in RTRs, half of the 18 positive warts also showed this suprabasal single-cell positivity, whereas the other half showed strong positive staining in a remarkable pattern of segmental positive suprabasal full epithelium thickness columns (zebroid pattern) (Figure 1B, D and Figure 2). This particular pattern was not linked to eccrine ducts or hair follicle structures; the latter actually seemed to be spared. This zebroid pattern correlated with retinoid treatment (topical and systemic) when comparing retinoid-treated RTRs (warts and in situ SCCs) with non–retinoid-treated RTRs (Table 2) (P < .001). Only 1 patient without (anamnestically traceable) retinoid treatment exhibited the same K13 expression pattern.

Most warts were superficially excised, with no perilesional skin available. In 1 retinoid-treated RTR, the perilesional skin showed K13 positivity comparable with the previously described zebroid pattern. All warts in both groups were negative for K19, with use of retinoids having no demonstrable effect on K19 expression (Figure 1C-D).

**COMMENT**

Data from numerous animal and in vitro studies with cultured human keratinocytes indicate that retinoids affect epidermal proliferation and differentiation. Retinoids proved to repress differentiation-specific keratins (K1/K10) and strikingly induced expression of K13 and K19, 2 keratins that are only coexpressed in fetal skin and are not usually present in the epidermis of adults. In contrast to these findings of coupled K13 and K19 induction by retinoids, Agarwal et al were the first to report uncoupled regulation of K13 and K19 expression in a human SCC cell line.

Our in vivo data with immunohistochemical evaluation of K13 and K19 expression in warts of RTRs vs ICIs show that retinoids used as chemoprotective agents in

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Table 1. K13-Positive and K13-Negative Warts in RTRs and ICIs

<table>
<thead>
<tr>
<th>Warts, No. (%)</th>
<th>RTRs (n = 21)</th>
<th>ICIs (n = 21)</th>
</tr>
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<tbody>
<tr>
<td>K13 positive</td>
<td>18 (86)</td>
<td>3 (14)</td>
</tr>
<tr>
<td>K13 negative</td>
<td>3 (14)</td>
<td>18 (86)</td>
</tr>
</tbody>
</table>

*K13 indicates keratin 13; RTRs, renal transplant recipients; and ICIs, immunocompetent individuals.*
RTRs for preventing skin cancer also strongly relate only to K13, and not K19, expression. Our finding of retinoid therapy–related uncoupled K13 and K19 expression in these patients could be 3-fold. First, the retinoid concentration in our patients could be sufficient to enhance K13 expression but not K19 expression. Earlier findings in human epidermal cultures showed stronger induction of K13 than K19 by retinoids with an already marked increase of K13 in response to low levels of retinoids, whereas for K19 induction a higher retinoid concentration was necessary. Although the dosage of systemic retinoids taken by many of our patients had to be lowered during treatment because of severe mucocutaneous adverse effects, no induction of K19 was found in patients still receiving the higher dosages of acitretin. Second, the response of keratinocytes to retinoids in vitro might not be representative of the response in vivo, and retinoids in humans in vivo might not induce an embryonic type of differentiation and might only selectively induce K13 expression. In the only previous in vivo study on the effects of retinoids in photo-aged skin only K13, and not K19, expression was studied. Finally, this differentiation toward esophageal-type epithelium in contrast to so-called embryonic-type differentiation found in animals and in vitro could be specific for skin and skin lesions in RTRs; earlier studies already showed that effects of retinoids differ in normal compared with diseased keratinocytes.

Retinoid treatment significantly correlated with a specific pattern of K13 expression in skin lesions of RTRs. This pattern, which we termed zebroid because of alternating suprabasal columns of K13-positive and K13-negative keratinocytes, is suggestive of a genetic mosaicism, reflecting clonal expansion of genetically altered stem cells. In warts and slightly dysplastic skin of RTRs, segments of epidermis may contain keratinocytes with a genetic abnormality, making them more susceptible to inductive actions of retinoids. Future studies might unravel the underlying genes that are involved in this process and whether, for instance, transforming types of human papillomavirus might play a role.

Interpretation of the biological impact of K13 expression in retinoid- and non–retinoid-treated warts of RTRs could be 2-fold. The first interpretation relates the presence of K13 in skin to differentiation, in parallel to internal squamous epithelia, in which K13 is restricted to the suprabasal epithelial compartment and is associated with differentiation. Retinoids, by inducing K13 expression or directing keratinocytes toward (esophageal) differentiation, might be chemopreventive by “freezing” cells in this differentiated state and preventing them from (further) dedifferentiating. Results of previous studies of retinoid effects on epidermal keratinocytes have shown that in response to retinoid treatment, higher molecular weight keratins, typically encountered in squamous epithelia, disappear, and synthesis of 2 new low molecular-weight keratins, a 40- and a 52-kd keratin, corresponding to K19 and K13, respectively, is enhanced. In the absence of vitamin A, the opposite occurs, with an enhanced terminal epidermal type of differentiation. Retinoid-induced esophageal-type differentiation could provide an explanation for the cosmetic improvement of lesional skin in these RTRs, who often had multiple hyperkeratoses before treatment: esophageal epithelium is, in contrast to the keratinizing epidermis, a nonkeratinizing squamous epithelium. By inducing nonkeratinizing differentiation, retinoids could lower the number of hyperkeratotic skin lesions. As an adverse effect, in normal skin the diminished cutaneous keratinization caused by retinoids leads to desquamation of palms and soles, which usually show the most prominent keratinization. On the lips, the outer cutaneous side becomes more vulnerable because of differentiation toward wet epithelium, leading to chelitis, another known adverse effect of acitretin treatment also present in our patients.

The second interpretation relates the presence of K13 to a more dedifferentiated and potentially malignant phenotype. Regarding cutaneous carcinogenesis, malignant transformation is heralded by a switch from production of high-molecular-weight keratins usually present in adult skin (K1/K10) to low-molecular-weight keratins also characteristic of fetal skin and simple epithelia (eg, K8/K18 and K19). Presence of K13, a low-molecular-weight embryonic keratin, would fit within this concept. It is tempting to attribute relevance to the high frequency of K13 in warts of RTRs and to speculate that it may be related to the higher susceptibility of warts in these patients to become malignant. This would be analogous to mouse models on skin carcinogenesis in which aberrant K13 expression is a consistent finding in chemically and v-Haras–induced papillomas and SCCs. In situ SCCs of RTRs and ICIs we found frequent K13 expression in 75% and 45% of lesions, respectively (20 in situ SCCs...
tested within each group, data not shown), which is in concert with this second hypothesis. The pattern of K13 expression in in situ SCCs of both groups was different from the retinoid therapy–related K13 expression (zebroid pattern, compare Figure 1B, G). Only 4 RTRs with in situ SCCs used retinoids, and in these patients the retinoid therapy–related zebroid K13 pattern was most pronounced or only present in perilesional, slightly dysplastic skin (Figure 1H).

When this latter interpretation would be applicable to retinoid-related K13 expression, use of retinoids might actually be dangerous for these patients. This is contradicted by studies of the long-term safety of retinoid therapy, since no increased incidence of skin cancer is reported.29 Studies of skin cancer chemoprophylaxis with retinoids in RTRs actually showed reduction in the skin cancer incidence.8

In conclusion, this retrospective in vivo study of embryonic keratin expression in warts of RTRs shows that retinoids strongly relate to K13 but not K19 expression. By keeping keratinocytes in this esophageal-type differentiation, retinoids might act chemopreventively. Retinoids correlate with a highly distinctive and strong K13 expression, which we termed zebroid, making K13 a useful marker for evaluating retinoid treatment in these patients. The alternating zebroid K13 pattern is suggestive of an underlying genetic mosaicism. Even in the absence of retinoids, a significantly higher percentage of K13 positivity is found in warts of RTRs compared with warts of ICRs. Future prospective, randomized, and well-controlled studies need to establish the relevance of this finding and whether K13, in analogue to mouse models of skin carcinogenesis, might become a predictive marker for malignant progression.

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