Keratosis lichenoides chronica (KLC) is a rare chronic progressive cutaneous disease that is part of the heterogeneous group of lichenoid dermatoses. The typical clinical presentation is characterized by lichenoid hyperkeratotic papules and nodules arranged in a linear and reticular pattern on the trunk and extremities. Our case confirms the existence of a vascular variant of KLC. There is no consensus about its treatment, since it is refractory to many different treatment modalities. We report the effectiveness of acitretin in KLC in combination with tacalcitol. KLC is of unknown aetiology, but is perhaps associated with systemic diseases, most importantly glomerulonephritis and lymphoma. This is the second case associated with hypothyroidism. **Key words: acitretin; hypothyroidism; keratosis lichenoides chronica; tacalcitol.**

There is presently no consensus about the treatment of KLC, since it is refractory to many different treatment modalities. We report here the effectiveness of acitretin in combination with tacalcitol.

**CASE REPORT**

A 48-year-old woman presented with a pruritic lichenoid papular eruption, the intensity of which had waxed and waned over a period of 20 years. She experienced an improvement in the summertime, especially when she was exposed to the sun in southern Europe. She described having had a seborrhiec-dermatosis-like eruption of KLC. Since the age of 15, she had been hyperactive and had been treated with a variety of hormonal contraceptives for 20 years. At the age of 42, she developed hypothyroidism and was treated with levothyroxine since 1997. The eruption was well-regulated during the time the patient was treated with levothyroxine. Since 1998, her hypothyroidism was treated with 60 μg levothyroxine and 25 μg liothyronine daily. At the age of 47, she developed a neurological disorder that was diagnosed as a transient ischemic attack. She was treated with a statin, which was continued after the attack. Since 2000, she has been on hormone replacement therapy. She is a non-smoker and is in good health.

Keratosis lichenoides chronica (KLC), a rare chronic progressive cutaneous disease that is part of the heterogeneous group of lichenoid dermatoses (1), was described by Kaposi more than a century ago as ‘lichen ruber acuminatus verrucosus et reticularis’. In 1938, Nekam reported a similar case and named it ‘porokeratosis striata lichenoides’ (2). Eventually, in 1972, Margolis gave the dermatosis its widely accepted name of KLC. Still, however, KLC remains a source of controversy, probably because of its rarity and the absence of a clear definition. At present it is accepted as a distinct entity, clinically and histologically, at one end of the spectrum of lichen planus (3). The typical clinical presentation of KLC is characterized by lichenoid hyperkeratotic papules and nodules arranged in a linear and reticular pattern on the trunk and extremities. Less frequently, seborrhiec-dermatitis-like lesions on the face, palmoplantar keratoderma, nail changes and mucosal lesions have been described (2, 4–7). A rare vascular variant of KLC has also been described (8). KLC is of unknown aetiology, but may be associated with systemic diseases, most importantly glomerulonephritis and lymphoma (2). One previous case was associated with hypothyroidism (9).
tion on the face a year previously. Furthermore, she experienced recurrent oral aphthosis. The patient’s medical history showed an auto-immune hypothyroidism since the age of 2 years, for which she was given thyroxine. She did not take any other drugs. Family history did not reveal significant diseases.

Physical examination showed symmetric clusters of erythematous lichenoid hyperkeratotic papules and several excoriations on the upper back, the shoulders, the lateral aspects of the trunk, and the buttocks (Fig. 1). The flexor side of the arms and forearms showed an extensive reticular network of telangiectasias bilaterally, with distinct linear hyperkeratotic ridges and papules (Fig. 2). There was no oral aphthosis or genital ulcers. The nails were not affected.

Histology of the lichenoid papules showed a superficial perivascular infiltrate forming a mild band-like infiltrate which consisted of lymphocytes and histiocytes, the latter often loaded with melanin pigment.

Moreover, we noticed multiple eosinophilic colloid bodies in the papillary dermis and an extensive vacuolar alteration of the basal keratinocytes. High in the spinous layer there were multiple necrotic keratinocytes. Furthermore, there was epidermal hyperplasia with several V-shaped spots of hypergranulosis, focal spongiosis and parakeratosis. On direct immunofluorescence the colloid bodies stained with IgM, and a focal immunoreactivity to fibrinogen was seen at the dermal-epidermal junction.

Routine laboratory examinations, including erythrocyte sedimentation rate, complete blood count and liver and kidney function tests were within normal limits. Immunologic investigation showed a positive ANA of 1/80 with fine granular staining pattern, and an elevated IgE (234 U/ml). The thyroid function was well controlled. Hepatitis C serology was negative.

The treatment consisted of tacalcitol once a day and acitretin, which was initiated at a dose of 10 mg a day. She tolerated the acitretin well, so the dose was increased to 30 mg a day and led to an almost complete clearing of the lichenoid eruption, leaving only a slight erythema, within 4 weeks. Also the telangiectasias appeared less prominent than before, but did not completely vanish on the inner side of her arms. After 4 months, acitretin was tapered back to 10 mg a day as maintenance therapy in association with tacalcitol, with success. During a 9-month follow-up, the improvement was maintained.

**DISCUSSION**

The great variety of synonyms used for KLC implies that there is no complete consensus about this rare disorder. Braun-Falco et al. (10), who reviewed 37 cases of KLC, confirmed the three hallmarks introduced by Pinol-Aguada, namely, lichenoid papules, linear hyperkeratotic ridges and erythematous plaques. Our patient experienced large clusters of itchy lichenoid hyperkeratotic papules on the back without a vascular background. On the inner aspects of the arms we noticed a prominent network of telangiectasias with reticular hyperkeratotic ridges and less prominent papules. We also noticed several other characteristic clinical features of KLC, namely: the history of sun-induced improvement, the seborrheic-dermatitis-like lesions on the face, and the recurrent aphthosis (2). Nail involvement, palmar-plantar hyperkeratosis, oral, genital and the eye involvement, are less frequently reported in patients with KLC (4–7). KLC is a polymorphous dermatosis, and there is little uniformity in the diagnostic criteria (2).

As regards the distinct vascular network on the arms we refer to cases reported in the French literature as ‘Keratose lichenoid striée’, the cases of Wätzig et al. and David et al., who described it as a vascular variant of KLC (8, 11–13). Noteworthy is the striking similarity of the telangiectatic pattern on the inner side of the arms with the latter report. Interestingly, both cases were associated with severe itching, whereas KLC is normally asymptomatic, suggesting a possible relation between the itch and this subtype of KLC. Microscopic examination of the biopsy specimen showed a lichenoid pattern, although the dense band-like infiltrate, which is so characteristic for this group, was only mild in our case. However, in 13% of cases the typical lichenoid infiltrate was absent in a comparative pathological study of KLC (2). In conclusion, our clinical and histological observations are compatible with KLC.

KLC has a progressive course that may wax and wane chronically, although there are two reports of spontaneous resolution after 7 and 13 years, respectively (14, 15). Unfortunately, it is refractory to various treatment modalities. Several topical ointments, such as potent steroids, tretinoin, anthralin and keratolytics have failed to give significant improvement (2). Recently, calcipotriol twice a day was reported to give good results in two patients with KLC (9, 16). Nevertheless, tacalcitol was proposed because of the once a day application. The largeness of the affected hyperkeratotic area obliged us to add a systemic therapy. An oral retinoid was started, since this drug interferes with the epidermal proliferation and has anti-inflammatory actions (17). Moreover, the other vascular variant of KLC responded well to etretinate, and recently Avermaete et al. reported a case successfully treated with acitretin (8, 18). We therefore started acitretin 10 mg a day and increased the dose to 30 mg a day (0.5 mg/kg/day), since she tolerated it well. This led to an almost complete clearing of the lichenoid eruption after 4 weeks and complete relief from the itching. The telangiectasia also appeared less prominent than before: we noticed a reduction in the redness and thickness of the cutaneous blood vessels. This was not the case in the other vascular variant of KLC (8). An alternative systemic treatment would have been PUVA, but the
data about its effectiveness are not consistent in the literature (2, 19–21). Systemic corticosteroids, methotrexate, cyclosporine, dapsone, vitamin A, gold, and tetracyclines are mentioned sporadically, but have been ineffective (2).

The patient presented by Grunwald et al. had had hypothyroidism for years, which was treated with thyroxine (9). Because our patient suffers also from hypothyroidism and is treated with thyroxine, we would like to draw attention to this association. Further cases are needed to evaluate a possible relationship. Concerning possible other associated diseases, two reports of chronic hepatitis related to keratosis lichenoides chronica – Wätzig & Schaarschmidt (13) and Marschalko et al. (22) – ought to be mentioned.

REFERENCES