over several weeks to months. Additionally, response to antibiotics has been described in HKFE cases without associated BAK exposure. For completeness, patch testing could have been considered to rule-out BAK allergic contact dermatitis; however, as the clinical features were characteristic for HKFE and our patients responded rapidly to initial therapy with amoxicillin-clavulanic acid, we did not pursue this.

This observation in a mother–daughter pair, while not sufficiently robust to claim genetic aetiology, raises the possibility of genetic factors contributing to HKFE/GP, as also suggested by Robinson et al. (2019). The microbiome and skin barrier function are genetically influenced and both are involved in HKFE pathogenesis.

The reversal of dysbiosis with antibiotics is an effective treatment strategy in conditions including erythrasma, confluent and reticulated papillomatosis, and also HKFE. Positive bacterial cultures have been occasionally reported in HKFE, such as in Case 1, with organisms including Klebsiella pneumoniae, Streptococcus milleri and Coagulase-negative Staphylococci. These organisms are normal skin commensals, and we believe the positive cultures reflect the state of dysbiosis rather than bacterial infection. However, further studies are required to characterise the altered microbiome of these patients, which may help determine the pathogenicity of organisms in HKFE.

Case Letter

Dear Editor,

Keratosis lichenoides chronica: A case report and focused overview of the literature

Keratosis lichenoides chronica (KLC) is an inflammatory skin disorder, characterised by diffuse, lichenoid hyperkeratotic papules arranged in an either linear or reticular fashion, erythematousquamous plaques and seborrheic-like dermatitis on the face. KLC runs a chronic, progressive course and demonstrates poor response to treatment. Herein, we report the case of a 55-year-old woman referred to our Dermatology Unit for a 4-year history of diffuse, persistent, slightly pruritic cutaneous lesions, predominantly affecting her face and limbs. Previously, she had been diagnosed with pityriasis lichenoides chronica and then with discoid lupus erythematosus. Topical corticosteroids, calcineurin inhibitors, phototherapy, hydroxychloroquine and methotrexate had been tried with little to no benefit. Physical examination revealed numerous, 2–4 mm diameter, keratotic, red-purplish papules symmetrically distributed to the upper and lower limbs and arranged in a linear-reticular pattern. On the face, there were well-demarcated erythematous papules and plaques with yellowish scales, especially in seborrhoeic areas. Cutaneous lesions were reported to improve with sunlight exposure. Concomitant blepharitis with a painful erosion on the tarsal conjunctiva of the left eye was an additional finding. Finally, a background photodistributed erythema was documented (Fig. 1). No involvement of scalp or oral

Figure 2: Response of Hyperkeratotic Flexural Erythema to Amoxicillin–Clavulanic Acid in Case 2. (a) Scaly, erythematos plaques in the right groin at time of onset. (b) 5 months later (at first presentation), progression of lesions to involve the bilateral groin and thighs. (c) Marked improvement of the lesions after 7 days of amoxicillin–clavulanic acid.

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and genital mucosae was recorded. Laboratory examinations were within normal ranges. Histopathological examination revealed focal epidermal parakeratosis and hypergranulosis, band-like lymphocytic infiltrates in the dermis and exocytosis of lymphocytes at the dermal–epidermal junction (Fig. 2). Direct immunofluorescence showed granular Immunoglobulin M (IgM) deposits at the dermal–epidermal junction and within dermal papillae. Based on clinical and histological features, a diagnosis of keratosis lichenoides chronica was made. The patient refused testing for nucleotide-binding domain and leucine-rich repeat containing proteins 1 (NLRP1) gene.

Also known as Nekam’s disease, KLC is an exceptionally rare dermatosis, with only 82 cases reported to date (Table S1). It shows a peak of incidence in the fourth decade, with a slight male predominance. About 25% of cases are paediatric. KLC is hallmarked by asymptomatic, violaceous, scaly papules arranged in a reticular pattern over the body, more commonly on the limbs, and showing either seborrhoeic dermatitis-like or rosacea-like features on the face.1 Palmoplantar keratoderma may also occur. Less frequently, oral and genital mucosae can be affected with ulcerative lesions, while ocular involvement usually manifests with blepharitis, conjunctivitis, uveitis or iridocyclitis. Cutaneous appendages can be involved as well, with onychodystrophy or alopecia.2 Histopathological findings typically include hyperkeratosis with focal parakeratosis, irregular acanthosis admixed with areas of atrophy, vacuolar degeneration of the basal layer and mixed inflammatory lichenoid infiltrates in the upper dermis (often around infundibula and acrosyringia).3 Further complicating its diagnosis, several clinicopathological variants of KLC have been described, including vascular, purpuric, lupus-like, porokeratotic associated to amyloidosis and generalised forms (Table S2).

Existing therapeutic options include systemic retinoids (acitretin, isotretinoin and etretinate) and phototherapy.

Figure 1 Typical KLC clinical features in a 55-year-old woman: (a) erythema and scaling in seborrheic areas; (b) red, keratotic papules arranged in a linear-reticular pattern; (c) a verrucous plaque on dorsal feet surface; (d) close up view of lesions localised on the right thigh.

Figure 2 Histopathological examination revealed presence of epidermal focal parakeratosis, hypergranulosis and lymphocytic band-like infiltrates in the dermis, with exocytosis of lymphocytes at the dermal–epidermal junction (Haematoxylin & Eosin, X20).
(psoralen-UVA, narrow band-UVB), which can lead to complete remission in almost half of patients, alone or in combination. Systemic steroids, antimalarials and antibiotics showed minimal effectiveness.²

The pathogenesis of KLC has not yet been elucidated fully. Traditionally, KLC was identified as a variant of LRP, and was long presumed to have the same underlying immune-mediated pathophysiology.³,⁴ Beyond clinical and histopathological differences, reclassification of KLC as a separate entity was recently substantiated by the discovery of a gain-of-function mutation in nucleotide-binding domain and leucine-rich repeat containing proteins 1 (NLRP1) gene in a family with semi-dominantly inherited KLC.⁵ NLRP1 is an inflammasome sensor protein highly expressed in keratinocytes and cutaneous fibroblasts. In response to a variety of stimuli, NLRP1 activates caspase-1 leading to increased production of interleukin (IL)-1 and IL-18. The KLC-causing mutation disrupts a leucin-rich repeat (LRR) domain, resulting in constitutive NLRP1 self-dimerisation and inflammasome activation.⁵ Such autoinflammatory setting determines reactive keratinocyte proliferation, which translates into parakeratotic hyperkeratosis on histology.¹,⁵ Over time, excessive keratinocyte turnover may favour the rise of neoplastic skin lesions, such as multiple keratoacanthomas.⁵,⁶

Accordingly, familiar KLC was included within the spectrum of autoinflammatory keratinisation diseases, an umbrella term encompassing monogenic skin diseases with an autoinflammatory pathogenesis.⁷ Germinal NLRP1 gene mutations have also been described in patients with dyskeratotic skin manifestations (e.g. follicular hyperkeratotic papules), polyarthritides and recurrent fever: suitably, this newly discovered disorder was named NLRP1-associated autoinflammation with arthritis and dyskeratosis (NAIAD).⁸ Such novel insights into familiar KLC may pave the way for anti-IL-1 therapeutic trials also in sporadic cases. Indeed, there are no reports on IL-1 receptor antagonist (i.e. anakinra) treatment in familiar KLC, although good response to anakinra was recorded in a case of NAIAD.⁸

Further research will be needed to better understand the pathophysiology of both familiar and sporadic KLC cases.

**ETHICS APPROVAL**

The study was conducted in accordance with the ethical standards of the responsible committee on human experimentation (institutional and national), with the Helsinki Declaration of 1975, as revised in 2000, and with the Taipei Declaration.

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**Data availability statement**

Anonymised data will be shared upon reasonable request from any qualified investigator for purposes of replicating procedures and results.

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**DETAILS OF CONTRIBUTION OF INDIVIDUAL AUTHORS**

Italo Francesco Aromolo, Serena Giacalone, Giovanni Genovese and Carlo Alberto Maronese equally participated in data acquisition, analysis, interpretation and drafting of the manuscript. Angelo Valerio Marzano participated in study concept and design and supervised the study. All authors critically revised the manuscript for important intellectual content and approved the final manuscript.

**PATIENT CONSENT STATEMENT**

Written informed consent was obtained from the patient included in the study, regarding also the publication of the photos.

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Dear editor,

A case of aggressive and protuberant nodular fasciitis on the finger with repeated contact

Nodular fascitis (NF) is a benign proliferation of myofibroblasts with a predilection for the head, neck, trunk and upper extremities of young adults. It originates commonly in the subcutis but may also arise in the dermis, fascia and skeletal muscle. Here, we report a case of NF on the index finger that developed rapidly and presented a protuberant appearance.

The patient was a 16-year-old boy who performed Kendo, a traditional martial arts form in Japan. He experienced intermittent irritation and had a callus-like eruption on the basal phalanx of the left palm index finger for a year. One month prior to his first visit, the surface of the eruption became ulcerated (Figure 1a), and during the initial examination, a dome-shaped erythematous mass measuring 16–18 mm in diameter was observed (Figure 1b). Suspecting a pyogenic granuloma (PG), we excised the nodule, which had grown rapidly to 20 mm in diameter within a week (Figure 1c, d).

Histopathological findings revealed an ulcerated tumour with low cell proliferation in the edematous outer layer and more cellular areas forming multiple irregular bundles as a storiform pattern in the inner layer (Figure 1e, f). In edematous lesions, spindle cells proliferate in the myxoelematous stroma accompanied by extravasation of erythrocytes and inflammatory cell infiltration (Figure 1g). Immunohistochemically, the tumour cells were positive for αSMA (Figure 1h) but negative for CD34 (Figure 1i), desmin, AE1/AE3, EMA or S-100 protein, suggesting a characteristic of a myofibroblastic tumour. CD68 staining was positive in infiltrating monocytes but negative in tumour cells (Figure 1j). Although the aggressive nature of the tumour suggested malignancy, the histopathological analysis showed hypercellular, storiform pattern and partial myxoid degeneration without tumour necrosis or nuclear atypia; thus, we diagnosed this case as NF.

This case showed an atypical clinical course of NF. In addition to our case, five other cases of aggressive NF have been reported previously (Table S1). The mean age of onset for these cases was 27 years. These cases were clinically diagnosed as sarcoma or PG, and surgical treatment was performed in all cases. Five out of six lesions were on

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**Figure 1** (a–d) Clinical presentation. (a) One month before the first visit an erythematous mass with ulceration was present on the left-hand index finger. (b) A dome-shaped elevated erythematous mass of 16–18 mm in diameter seen on the left-hand index finger. (c) At the time of surgery, the mass had increased in size to over 20 mm in diameter. (d) Lateral view at the time of the surgery. (e–g) Histopathological findings with haematoxylin and eosin (H&E) staining. (e) Dense proliferation of atypical spindle-shaped cells in the dermis. (f) Higher magnification of the lesion consisting of multiple irregular bundles of cells in a storiform pattern in the inner layer. (g) Spindle cells are proliferating in the myxoelematous stroma (tissue culture like appearance) accompanied by erythrocyte extravasation and inflammatory cell infiltration in the outer layer. (h–j) Immunohistochemical findings. (h) Tumour cells were positive for αSMA. (i) Tumour cells were negative for CD34. (j) CD68 staining was positive in infiltrating monocytes and negative in tumour cells. Scale bar (e) 5 mm; (f–j) 200 μm.

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Informed consent: Informed consent has been obtained from the patient.

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