Nailfold capillaroscopic changes in Kindler syndrome

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Summary

Kindler syndrome (KS), the fourth major type of hereditary epidermolysis bullosa (HEB), is a rare, autosomal recessive disorder characterized by skin fragility and blistering at birth followed by development of marked photosensitivity and progressive poikilodermatous skin changes in later years. We reported here the case of a 54-year-old woman, who fulfills the diagnostic criteria of KS type of HEB, putting accent on the nailfold capillaroscopic changes. Using videocapillaroscopy we observed pronounced alterations in finger nail capillaries including reduction in capillary density, features of neoangiogenesis (architectural derangement, elongated loops, extremely tortuous, bushy or branching capillaries, thin, branching and interconnected capillaries), enlarged and giant capillaries. We consider the changes observed as an adaptive mechanism that compensate the loss of capillaries due to chronic periungual trauma. Further studies with larger number of patients are needed to confirm the significance of capillaroscopy findings for patients with HEB.

Keywords: Kindler syndrome, capillaroscopy

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micron) and giant capillaries (width > 50 micron); distribution: architectural derangement and features of neoangiogenesis; blood flow: normal; absence of hemorrhages and thromboses. The interpretation was: presence of nailfold capillary microangiopathy (Figure 2).

KS is caused by a deficiency of the protein kindlin-1 due to a mutation in the gene KIND1 mapped to chromosome 20p12. It manifests during the neonatal period and infancy with skin fragility and trauma-induced skin blistering, which usually resolve with age. Later, varying degrees of photosensitivity and progressive poikiloderma develop. Face, hands and feet are most affected. Ocular and nail changes, periodontitis, esophageal, anal, vaginal and urethral stenosis have been also observed. Additionally, KS patients present an increased susceptibility for development of squamous cell carcinomas. The diagnosis is essentially clinical but could be supported by histopathologic examination, immunostaining with anti-kindlin-1 antibody, electron microscopy examination, and molecular genetic testing (1-4).

In 2005, Angelova-Fischer et al. (5) proposed the following diagnostic criteria for the syndrome: i) Major criteria: acral blistering in infancy and childhood, progressive poikiloderma, skin atrophy, abnormal photosensitivity, gingival fragility and/or swelling; ii) Minor criteria: syndactyly and mucosal involvement (anal, esophageal, urethral, laryngeal stenosis); iii) Associated findings: nail dystrophy, ectropion, palmoplantar keratoderma, leukoplakia, squamous cell carcinomas, skeletal abnormalities, periodontitis and tooth decay. The presence of 4 major criteria makes the diagnosis of KS certain. The presence of 3 major and 2 minor criteria makes the diagnosis probable and the presence of 2 major criteria and 2 minor criteria or associated symptoms renders the diagnosis likely. Our patient had all 5 major criteria, and mucosal and nail involvement, ectropion, squamous cell carcinoma, periodontitis and poorly preserved teeth.

Figure 1. Ectropion (a); periodontitis (b); erythema, atrophy, hyper- and hypopigmented macules, telangiectasia, xeroderma, atrophic scars (b-e), atrophic scarring with shiny cigarette paper-like wrinkling and small ulcerations on dorsum of the hands (f).

Figure 2. Capillaroscopic findings: reduction in capillary density, shape heterogeneity, marked tortuosity with varied appearance, enlarged and giant capillaries, architectural derangement, neoangiogenesis. Magnification: ×60 (a); ×200 (b-d); ×500 (e-h).
So far there are not studies on microcirculation in HEB and KS in particular. That is why we aimed to evaluate the changes of nail fold microcirculation using non-invasive, digital videocapillaroscopy. We observed pronounced alterations in finger nail capillaries with the main feature of capillary neoformation including elongated loops, extremely tortuous, bushy or branching capillaries, thin, branching and interconnected capillaries. The presence of enlarged and giant capillaries makes the differentiation with Raynaud phenomenon somewhat difficult. Moreover, the clinical appearance of the patient’s skin of the hands impress on general practitioners and rheumatologists that she suffer from autoimmune connective tissue disease. However, the patient did not report any vasospastic episodes of her fingers provoked by cold and the repeatedly immunological tests were normal. We consider the neoangiogenesis observed as an adaptive mechanism that compensate the loss of capillaries due to chronic periungual trauma. The role of sensory polyneuropathy could be additionally discussed. However, it is probably a consequence of the general disorder, too.

To the best of our knowledge, this is the first report of microcirculatory alterations in KS. Further studies with larger number of patients are needed to confirm the significance of capillaroscopy findings for patients with HEB.

References


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