Kindler Syndrome with Severe Intestinal Involvement: A 31-year Follow-up

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Kindler syndrome (KS) is an autosomal recessive genodermatosis characterised by trauma-induced blistering, poikiloderma, mucosal inflammation and varying degrees of photosensitivity (1, 2). It is caused by mutations in FERMT1 (also known as KIND1), the gene encoding Kindlin1 (3, 4). More than 50 mutations have been reported to the Human Gene Mutation Database (HGMD® Professional Release 2012.4). In addition to the characteristic cutaneous features, KS patients occasionally have intestinal involvement, including diarrhoea or colitis, mainly in childhood (5, 6). However, severe intestinal complications of multiple ileal stenoses have seldom been described in KS patients, and it has remained unclear whether these are characteristics of KS or are coincidental.

To further clarify the phenotype, genotype and FERMT1-expression of KS with severe intestinal involvement, we thoroughly evaluated a patient who has been carefully studied by dermatologists at a single institution for 31 years.

CASE REPORT

Clinical findings. The 31 year-old male patient presented multiple blisters and erosions at birth, which at that time were diagnosed as dystrophic epidermolysis bullosa. Both his non-consanguineous parents and sister are healthy. Because of the patient’s marked blisters, he was referred to Sapporo City General Hospital at 6 days old and has been carefully followed up by dermatologists at the hospital for 31 years. He had blistering after mild trauma, progressive poikiloderma and marked atrophy on the dorsal aspect of the hands and feet, with a cigarette paper-like wrinkled appearance (Fig 1a, b). He has undergone repetitive dilatation procedures for anal, urethral and oesophageal stenosis since age 5. He was first diagnosed with KS at age 18, based on characteristic clinical features as well as negative COL7A1 mutation, as described previously (7). He now shows marked atrophy and progressive poikiloderma of the entire skin, several erosions on the buttocks and legs, and microstomia, all of which are typical KS cutaneous features (Fig. 1c, d). Since the age of 16 years, he has suffered from stomachaches after meals due to severe stenosis and ulcers in the intestines, which have met the criteria for Crohn’s disease. Treatment with prednisolone, methotrexate, mesalazine, azathioprine and infliximab had little effect; therefore, he underwent emergent partial ileotomy due to bowel perforation at the ages of 25, 26 and 29 years (Fig. S1a1). However, some strictures still cause obstruction, predominantly in the ileum, and they easily recur after repeated balloon dilatation. He has undergone balloon dilatation for ileal stenosis more than 10 times in the past 5 years. Recent colonoscopy showed ulcers and several strictures, mainly in the ileum. The mucosa of the colon and the rectum also showed erosions but no strictures. Histopathological examination of ileal specimens showed apparent epithelial detachment of the mucosa and inflammatory cell infiltrates in the lamina propria (Fig. S1b). Two skin biopsy specimens taken at the ages of 18 and 31 years were compared. Indirect immunofluorescence was performed by using monoclonal antibodies against type IV, VII and XVII collagen, α6 and β4 integrin, BP230, plectin, uncein and laminin-332. Type VII collagen and type IV collagen showed a broad reticulate labelling pattern in both specimens, whereas other antibodies showed a normal bright linear pattern, which are compatible with KS (Fig. 1e). Electron microscopy revealed reduplication of the basement membrane in both specimens, but there was not a significant difference between them (not shown).

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Fig. 1. A blister on the hand at birth (a). Atrophy of the hand at 5 years of age (b). Poikilodermatous change and erosion on the dorsum of the hand at 30 years of age (c). Poikilodermatous change on the back at 31 years of age (d). Broad, reticulate labelling pattern is seen with LH7.2 monoclonal antibodies against type VII collagen (e).
In this study, we identified the novel mutation c.1564 delC in exon 12 of the FERMT1 gene in the maternal allele, which is predicted to lead to nonsense-mediated mRNA decay. FERMT1 mRNA was undetectable by RT-PCR, although the paternal mutation cannot be identified. Despite the novel mutation, the cutaneous findings do not differ from the cases with other mutations. The literature review was unable to reveal any relationship between the FERMT1 locus (20p12.3) and loci of Crohn’s disease.

In conclusion, long-term follow-up of KS patient revealed that severe late-onset ileal strictures can also occur as one of the clinical features of KS. It is suggested that FERMT1 expression is disrupted by maternal c.1564 delC and unknown paternal null-mutations in the FERMT1 gene.

**REFERENCES**