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Brief Report

Dermatology

Spontaneous gene repair, also called revertant mosaicism, has been documented in several genetic disorders involving organs that undergo self-regeneration, including the skin. Genetic reversion may occur through different mechanisms, and in a single individual, the mutation can be repaired in various ways. Here we describe a disseminated pattern of revertant mosaicism observed in 6 patients with Kindler syndrome (KS), a genodermatosis caused by loss of kindlin-1 (encoded by *FERMT1*) and clinically characterized by patchy skin pigmentation and atrophy. All patients presented duplication mutations (c.456dupA and c.676dupC) in *FERMT1*, and slipped mispairing in direct nucleotide repeats was identified as the reversion mechanism in all investigated revertant skin spots. The sequence around the mutations demonstrated high propensity to mutations, favoring both microinsertions and microdeletions. Additionally, in some revertant patches, mitotic recombination generated areas with homozygous normal keratinocytes. Restoration of kindlin-1 expression led to clinically and structurally normal skin. Since loss of kindlin-1 severely impairs keratinocyte proliferation, we predict that revertant cells have a selective advantage that allows their clonal expansion and, consequently, the improvement of the skin condition.

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Revertant mosaicism in a human skin fragility disorder results from slipped mispairing and mitotic recombination

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Spontaneous gene repair, also called revertant mosaicism, has been documented in several genetic disorders involving organs that undergo self-regeneration, including the skin. Genetic reversion may occur through different mechanisms, and in a single individual, the mutation can be repaired in various ways. Here we describe a disseminated pattern of revertant mosaicism observed in 6 patients with Kindler syndrome (KS), a genodermatosis caused by loss of kindlin-1 (encoded by *FERMT1*) and clinically characterized by patchy skin pigmentation and atrophy. All patients presented duplication mutations (c.456dupA and c.676dupC) in *FERMT1*, and slipped mispairing in direct nucleotide repeats was identified as the reversion mechanism in all investigated revertant skin spots. The sequence around the mutations demonstrated high propensity to mutations, favoring both microinsertions and microdeletions. Additionally, in some revertant patches, mitotic recombination generated areas with homozygous normal keratinocytes. Restoration of kindlin-1 expression led to clinically and structurally normal skin. Since loss of kindlin-1 severely impairs keratinocyte proliferation, we predict that revertant cells have a selective advantage that allows their clonal expansion and, consequently, the improvement of the skin condition.

Introduction

Spontaneous gene repair, also called revertant mosaicism, has been observed in genetic disorders primarily involving organs that undergo self-regeneration, such as bone marrow, liver, and skin (1). Depending on the frequency of the reversion and the clonal expansion of the reverted cells, the phenotype may improve in a sense of natural healing (2). Genetic reversion can occur through recombination, back, or second-site mutations, and in a single individual, the disease-causing mutation can be mended in various ways (3). The accessibility of the skin facilitates the recognition of revertant mosaicism, which has previously been described in epidermolysis bullosa (4, 5) and ichthyosis (6).

Kindler syndrome (KS; OMIM 173650) is an autosomal-recessive skin fragility disorder that results from loss-of-function mutations in *FERMT1*, which encodes kindlin-1, a key regulator of integrin activation. KS is characterized by congenital skin blistering and photosensitivity, evolving to poikiloderma with pronounced skin atrophy (7). Poikiloderma refers to a combination of hypo- and hyperpigmentation, atrophy, and telangiectasias and is morphologically characterized by thin epidermis without rete ridges, focal vacuolization of basal keratinocytes, and pigment incontinence. Typically, KS skin shows multiple planes of cleavage at the dermal-epidermal junction (DEJ) and reduplication and interruption of the epidermal basement membrane. Consistent with the epidermal atrophy, basal keratinocytes exhibit minimal proliferation (8, 9).

Here, we describe 6 patients with KS caused by inherited *FERMT1* duplicating insertions and a disseminated pattern of revertant skin patches. The reversion mechanism in each investigated skin

patch was back mutation by slipped mispairing in direct nucleotide repeats. In addition, mitotic recombination was documented in some reverted areas.

Results and Discussion

Patient 1 (P1), who exhibited the hallmark feature of KS (poikiloderma with pronounced skin atrophy), was the index case, with innumerable normal-appearing skin patches. Subsequently, we observed a similar pattern in P2. Notably, both patients had *FERMT1* duplication insertion mutations, which led to frame shifts, premature termination codons, and loss of full-length kindlin-1, as demonstrated by immunoblotting of keratinocyte lysates (10). A systematic review of 24 additional KS patients revealed that all individuals harboring duplicating insertions (P3–P6; Table 1) exhibited the same skin pattern (Supplemental Figures 1 and 2; supplemental material available online with this article; doi:10.1172/JCI61976DS1). P1 and P2 were available for systematic investigation.

P1, a 29-year-old male born to consanguineous parents, was homozygous for c.456dupA (11). During a 3-year follow-up, we noted normal-appearing patches with preserved texture, measuring between several mm² and 15 cm², disseminated over the entire integument, that contrasted the atrophic, dry, hyperpigmented, and often erythematous skin (Figure 1, A–C). Immunofluorescence staining of a skin sample obtained from an atrophic area (P1-1) revealed absence of kindlin-1 at the DEJ (Figure 2B). In contrast, 3 normal-appearing patches (P1-2, P1-3, and P1-4) showed positive kindlin-1 signals over the entire length of the biopsy, similar to normal skin (Supplemental Table 1).

P2, a 24-year-old female homozygous for c.676dupC, exhibited multiple normal-appearing patches, measuring 0.5–2 cm², that were clearly distinguishable from the atrophic integument

Conflict of interest: The authors have declared that no conflict of interest exists.

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Table 1
Clinical and genetic features

Pt	Sex	Age (yr) ^A	Origin	Clinical features	Revertant patch characteristics			<i>FERMT1</i> mutations
					Number	Surface area	Distribution	
P1	Male ^B	29	Germany	Poikiloderma with marked atrophy, erythema, hypertrophic gingivitis, cheilitis angularis, urethral strictures, ectropion	Innumerable	Several mm ² to 15 cm ²	Entire integument	c.456dupA homozygous, p.Asp153ArgfsX4
P2	Female ^C	24	Kosovo	Poikiloderma with marked atrophy, hypertrophic gingivitis, esophageal stenosis, ectropion	<10	0.5 cm ² to 2 cm ²	Hands, lower legs	c.676dupC homozygous, p.Gln226ProfsX17 ^D
P3	Female ^C	17	Turkey	Poikiloderma with marked atrophy, photosensitivity, gingivitis	<10	0.5 cm ² to 3 cm ²	Hands	c.676dupC homozygous, p.Gln226ProfsX17 ^D
P4	Female ^C	21	Kosovo	Poikiloderma with marked atrophy, webbing of fingers, microstoma, esophageal stenosis, genital involvement	<10	0.5 cm ² to 3 cm ²	Hands, neck, legs	c.676dupC homozygous, p.Gln226ProfsX17 ^D
P5	Female ^C	11	Kosovo	Discrete poikiloderma and atrophy, colitis	>10	0.5 cm ² to 2 cm ²	Hands, lower legs	c.676dupC homozygous, p.Gln226ProfsX17 ^D
P6	Male ^C	9	Serbia/Greece	Discrete poikiloderma and atrophy, urethral strictures	Innumerable	Several mm ² to 1 cm ²	Hands, arms, legs	c.[676dupC];[1677G>A] ^E , p.[Gln226ProfsX17] ^D ;[Trp559X]

^AAge at last examination. ^BConsanguineous parents, not available for analysis. ^CNonconsanguineous parents, all heterozygous carriers of the mutations.

^DIdentical *FERMT1* SNPs. ^ETo our knowledge, c.1677G>A was not previously reported.

(Figure 1, D and E). Kindlin-1 immunofluorescence was negative in the affected skin, but positive (comparable to control skin) in 2 normal-appearing patches (Figure 2, D and E, and Supplemental Table 1).

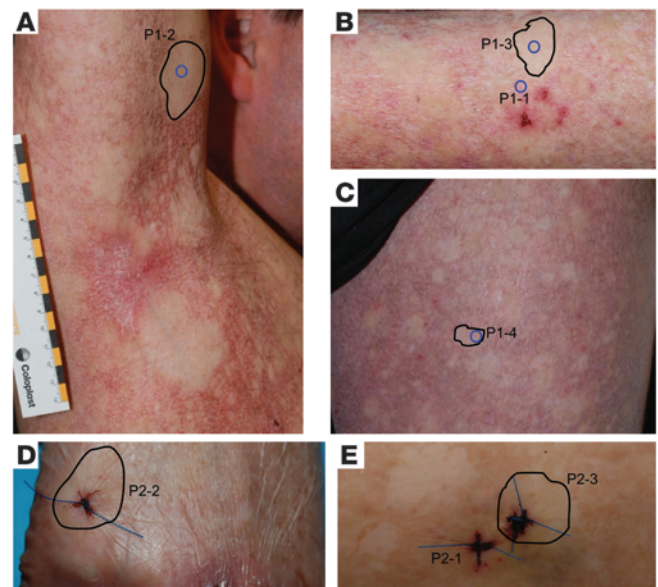
In both P1 and P2, restored kindlin-1 expression visibly improved skin morphology (i.e., preservation of rete ridges, epidermal thickness, and keratinocyte proliferation; Figure 2, C and E). Collagen VII staining was used to evaluate the integrity of the DEJ. In normal-appearing areas with positive kindlin-1 immunostaining, a linear collagen VII signal was seen that contrasted the irregular pattern in affected areas (Figure 2, B–E), which indicated that restoration of kindlin-1 normalized the DEJ.

These results strongly supported the hypothesis that the normal-appearing kindlin-1-positive spots represent areas in which the inherited mutations had reverted. To test this, we used laser dissection microscopy (LDM) to collect keratinocytes from areas with altered or normal DEJ. DNA was extracted, and *FERMT1* exons 4 and 5 were sequenced in P1 and P2, respectively. In both patients, the mutations were found in a heterozygous state in keratinocytes derived from all normal-appearing patches, whereas in diseased skin, the mutations were present in a homozygous state, similar to lymphocytes (Figure 3 and Supplemental Table 1).

Figure 1

Clinical features indicative of revertant mosaicism in KS. (A–C) P1 exhibited numerous normal-appearing skin patches of several mm² to 15 cm² on the entire integument, which remained constant over the 3-year observation period. Shown are the right axilla, upper thorax, and right arm (A); the left arm (B); and the left leg (C). (D and E) P2 had normal-appearing skin patches on the right hand (D) and the right lower leg (E). Revertant areas investigated here are outlined in black; biopsy sites are circled in blue or directly visible.

In the dermis below the reverted areas, the mutations were present in a homozygous state. Thus, the reversion had occurred only in keratinocytes, which are the sole kindlin-1-expressing cells in the skin (8, 9). The results were confirmed in P2 at the RNA level (data not shown), but no material was available from P1. Notably, in both patients, the same reversion mechanism (i.e., a single nucleotide deletion restoring the number of bases in the direct repeats) was found in all revertant patches, suggestive of a high propensity of the sequence to mutational events through slipped mispairing (12).



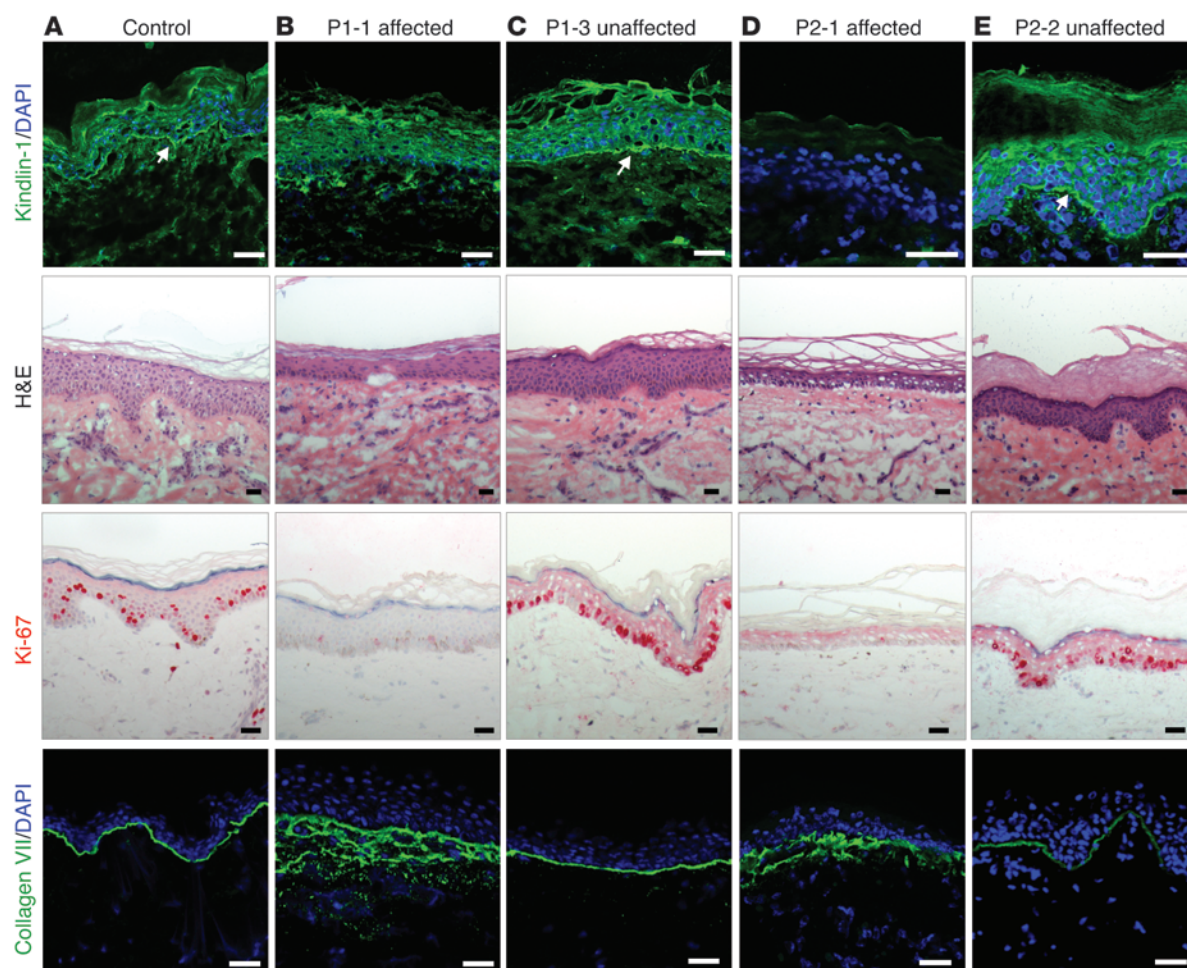


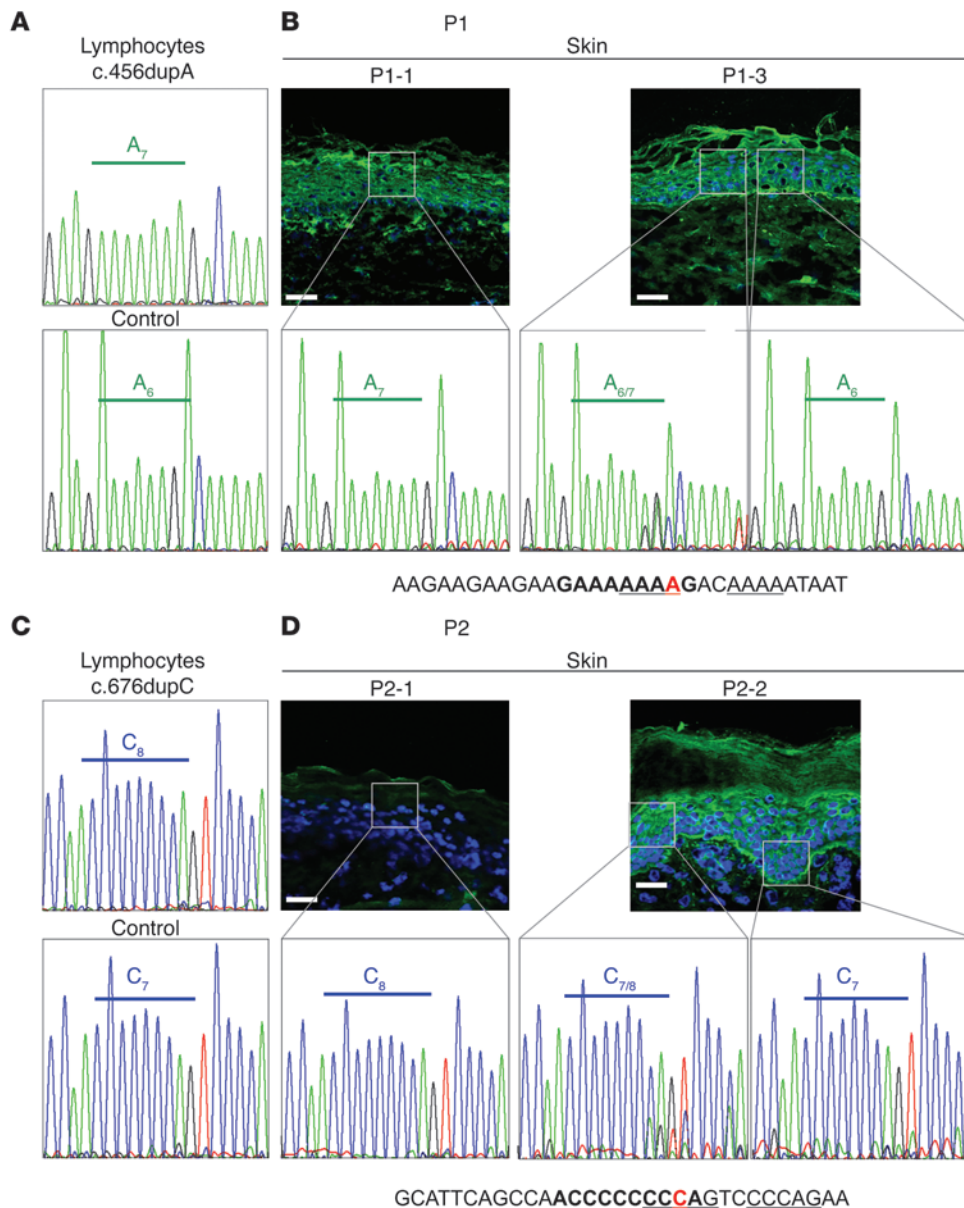
Figure 2

Kindlin-1 expression and skin morphology correlate with clinical phenotype. Skin of a control individual (**A**) and affected (**B** and **D**) and unaffected (**C** and **E**) skin of P1 and P2 were stained with H&E and antibodies to kindlin-1, Ki-67, and collagen VII. In the affected skin P1-1 and P2-1, loss of kindlin-1 expression was associated with thin epidermis, microblisters at the DEJ, strong reduction of Ki-67–positive keratinocytes, and irregular collagen VII staining at the DEJ and in the upper dermis. In contrast, in the normal-appearing patches P1-3 and P2-2 with restored kindlin-1 expression (arrows), skin morphology, Ki-67 staining, and collagen VII distribution were comparable to those of the control. Scale bars: 50 μ m.

Surprisingly, in both P1 and P2, the homozygous wild-type sequence was also disclosed in some areas of the normal-appearing skin samples (Figure 3), suggestive of a second mutational event in the revertant heterozygous keratinocytes — either a second back mutation, or mitotic recombination. To investigate this, we genotyped SNPs around *FERMT1* in DNA extracted from lymphocytes of P1 and P2 and identified the borders of the homozygous regions, which spanned about 1.1 and 1.4 Mb, respectively. In the revertant patches with the homozygous wild-type *FERMT1* sequences, recombination was confirmed by loss of heterozygosity of the SNP rs261360, located outside these homozygous regions (Supplemental Table 2).

These results demonstrated restoration of the *FERMT1* reading frame, kindlin-1 expression, and skin structure in the revertant patches. Revertant keratinocytes proliferated normally, having a selective advantage compared with kindlin-1–negative cells (Figure 2), allowing them to expand. Nevertheless, large reverted areas (~15 cm²) may represent confluence of multiple overlapping or neighboring revertant events.

The slipped mispairing model involves the misalignment of short direct repeats during DNA replication and represents a common mutational mechanism (13–15), which has previously been elucidated in microorganisms (16–18). The primer strand containing the newly synthesized repeat dissociates from the template strand and misaligns, leading to either insertion or deletion of 1 or more direct repeats in the continued DNA synthesis. The fidelity of DNA replication is dependent on the local sequence environment and the DNA polymerase (19). PolyA and polyC repeats, as observed in our patients, are known hotspots for microdeletions and microinsertions (20, 21). Prior meta-analysis of the sequence vicinity of a large number of microdeletions and microinsertions in different genes revealed that short oligonucleotide motifs significantly correlate with the occurrence of both mutational events (12). Such motifs — GA₆ and A₆G, or AC₆ and C₆A — were present close to the mutations in our patients (Figure 3). Additionally, the DNA polymerase β frameshift hotspot AAAAA, and the topoisomerase II consensus cleavage site CCCAG, which are potentially involved in site-specific recombination and putative deletion/insertion hot-

**Figure 3**

Identification of the reversion mechanism. LDM was used to collect epidermal keratinocytes from areas with altered or normal DEJ morphology. **(A)** Partial sequence of *FERMT1* exon 4 in lymphocytes of P1 showed the duplicating insertion in adenine repeats c.456dupA (A_7). The control had the adenine repeat sequence A_6 . **(B)** In affected skin sample P1-1, the mutation was found in a homozygous state. Analysis of DNA from microdissected areas (insets) of P1-3 from unaffected skin revealed the mutation in a heterozygous state ($A_{6/7}$) in the majority of the experiments, but the wild-type sequence (A_6) was also found. The sequence in the vicinity of the revertant mutation is shown below; bold denotes the GA_6-A_6G motif, underline denotes a DNA-polymerase β frameshift hotspot; red denotes the mutation. **(C)** Partial sequence of *FERMT1* exon 5 in lymphocytes of P2 and a control showing the duplicating insertion in a cytosine tract (C_8). **(D)** The homozygous duplication c.676dupC was disclosed in the affected skin P2-1, whereas analysis of DNA from microdissected areas (insets) in unaffected skin P2-2 disclosed the mutation in a heterozygous state ($C_{7/8}$), or the normal sequence (C_7). The sequence in the vicinity of the revertant mutation is shown below; bold denotes the AC_6-C_6A motif, underline denotes the vertebrate topoisomerase II consensus cleavage site; red denotes the mutation. Scale bars: 50 μ m.

spots, were close to the mutations. Based on the identical molecular event in all analyzed spots and the finding that of our cohort of 26, all 6 KS patients with duplicating insertions exhibited multifocal revertant mosaicism, we strongly favor this mechanism as the cause of reversion. A recent report of an 8-year-old boy homozygous for c.676dupC and having a normal-appearing skin patch on his hand (5) strengthens our findings. Recognition of multiple patches in young children is difficult (22), since small normal-appearing areas may only become visible with age, against the progressive poikiloderma (ref. 7 and Supplemental Figure 2).

The multifocal mosaic pattern in KS is similar to the rare disorder ichthyosis with confetti, which is caused by dominant *KRT10* mutations. In this disorder, multiple normal-appearing patches represent revertant areas in which the inherited mutation is rescued by different mitotic recombination events (6). In our cases, recombination occurred as a secondary mutational event, after the back mutation had already reverted the recessive condition.

These findings may open therapeutic perspectives for KS and other genodermatoses. The advantages of a therapy using revertant keratinocytes are the natural correction of the mutation and the lack of immune response. Grafting revertant keratinocytes in junctional epidermolysis bullosa demonstrated that the procedure is easy and well-tolerated by the patient, but cultivation of the reverted cells must be improved before it becomes clinically applicable (23).

Methods

Samples. EDTA blood was obtained from patients and parents. 4-mm punch skin biopsies were obtained from P1 and P2 (Supplemental Table 1 and Figure 1).

Immunomorphological analysis. H&E, immunohistochemistry, and indirect immunofluorescence staining of the skin was performed as described previously (8), using primary antibodies to kindlin-1 (8), Ki-67 (MIB-1; Dako), and collagen VII (LH7.2; Millipore) and the AEC (3-amino-9-ethyl-carbazole) system (Dako) or Alexa Fluor 488-conjugated goat anti-rabbit



and -mouse IgGs (Invitrogen). Nuclei were visualized with hematoxylin or DAPI. Stained sections were observed with light microscopy (Nikon 80i) or confocal laser scanning microscopy (LSM510; Carl Zeiss).

Mutation detection. Mutation detection on DNA extracted from EDTA blood (Qiagen kit) was performed as described previously (24). All 15 exons and adjacent junctions of *FERMT1* were amplified and sequenced in an ABI 3130XL genetic analyzer using Big Dye Terminator Chemistry (Applied Biosystems). DNA sequences were compared with the NCBI reference (NC_000020.10) using Mutation Surveyor DNA (2.61 Softgenetics). Mutations were verified by sequencing in both directions from independent PCRs. Additionally, 34 SNPs covering 7 Mb around *FERMT1* were genotyped by direct sequencing from the patients' lymphocytes (Supplemental Table 2).

Nested PCR was used to detect mutations in DNA from microdissected tissue (Supplemental Table 3). All PCRs were repeated with templates from at least 3 separate DNA isolations obtained by LDM, and all products were sequenced in both directions.

DNA and RNA isolation from skin sections. For DNA recovery by LDM, 5- μ m skin cryosections were mounted on 1.0-mm PEN membrane-covered slides (Zeiss). Collagen VII staining was used to visualize the DEJ because the signal is strong enough not to be bleached under LDM. Approximately 200 cells from affected and reverted areas were dissected with the Laser Robot Microbeam System (P.A.L.M. Microlaser Technology AG) and separately collected in the caps of 0.5-ml thin-walled tubes (Zeiss). For proteinase K digestion, 30 μ l of a mix consisting of PCR buffer (JumpStart RED Taq DNA Polymerase), water, and proteinase K (Qiagen) was added, and the tubes remained inverted for 60 minutes at 55 °C. Inactivation was subsequently achieved by heating to 98 °C for 15 minutes (2). For RNA isolation, 8 10- μ m skin sections were extracted with the Qiagen FFPE RNA kit (Qiagen). Reverse transcription was performed using the Advantage

RT-for-PCR Kit (Clontech) with 0.2 μ g total RNA, using oligodT primers. Primers were designed with Primer3 (<http://frodo.wi.mit.edu/primer3>; Supplemental Table 3).

Cloning. The entire epidermis was dissected with LDM from biopsies P1-2 and P2-3. Subsequently, DNA was isolated and submitted to nested PCR, and the amplicons were subcloned into the TOPO TA-cloning vector (Invitrogen). For each reaction, 15–20 clones were sequenced using the M13-RV primer.

Study approval. The study was approved by the Ethics Committee of the University of Freiburg and conducted according to the Declaration of Helsinki. Patients provided informed consent prior to their participation.

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