Revertant mosaicism in junctional epidermolysis bullosa due to multiple correcting second-site mutations in LAMB3

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Revertant mosaicism due to in vivo reversion of an inherited mutation has been described in the genetic skin disease epidermolysis bullosa (EB) for the genes KRT14 and COL17A1. Here we demonstrate the presence of multiple second-site mutations, all correcting the germline mosaicism. Both probands had a severe reduction in laminin-332 expression in their affected skin. Remarkably, the skin on the lower leg of patient 078-01 (c.628G→A/c.1903C→T) became progressively clinically healthy, with normal expression of laminin-332 on previously affected skin. In the other proband, 029-01 (c.628G→A/c.628G→A), the revertant patches were located at his arms, shoulder, and chest. DNA analysis showed different second-site mutations in revertant keratinocytes of distinct biopsy specimens (c.565-3T→C, c.596G→C;p.G199A, c.619A→C;p.K207Q, c.628+42G→A, and c.629-1G→A), implying that there is no a single preferred mechanism for the correction of a specific mutation. Our data offer prospects for EB treatment in particular cases, since revertant mosaicism seems to occur at a higher frequency than expected. This opens the possibility of applying revertant cell therapy in mosaic EB of the LAMB3 gene by using autologous naturally corrected keratinocytes, thereby bypassing the recombinant gene correction phase.

Introduction

Reverse mutations in germline or somatic cells bearing an inherited disease-causing mutation can change the phenotype from affected to normal by reexpression of the involved protein. These reverse mutations can be true back mutations — leading to the original wild-type sequence and thus wild-type protein — or can be additional second-site mutations that compensate for the effects of the primary inherited mutation (1). In the latter case, small changes in the amino acid sequence may occur. Revertant mosaicism has been demonstrated for different genetic diseases and in different cell types, as in hepatocytes, lymphocytes, and, in the case of epidermolysis bullosa (EB), keratinocytes (reviewed in refs. 2, 3). EB is a clinically heterogeneous group of heritable blistering disorders leading to fragility of the skin and mucous membranes. The subgroup junctional EB (JEB), characterized by separation at the lamina lucida of the epidermal basement membrane zone (BMZ), is caused by recessive mutations in the genes encoding type XVII collagen (COL17A1), integrin α6β4 (ITGA6 and ITGB4), or laminin-332 (LM-332; LAMA3, LAMB3, and LAMC2) (4). Recently, the first instance of correction of EB skin by transplantation of genetically modified epidermal stem cells was reported for a compound heterozygous carrier of laminin β3 (LAMB3) mutations (5).

Revertant mosaicism in EB due to in vivo reversion of somatic cells has been described for the genes COL17A1 and KRT14 (6–10). The first reported case involved the reversion of one of the defective COL17A1 alleles into a wild-type sequence due to a mitotic gene conversion event (6). In a second EB patient, the focal expression of type XVII collagen in the skin was the result of a second-site frame-restoring COL17A1 mutation (7). Recently, we demonstrated the occurrence of multiple correcting COL17A1 mutations in distinct type XVII collagen-positive skin patches in 2 unrelated probands (10). Amelioration of the EB phenotype can also be achieved on the RNA level by exon skipping, although this rescuing mechanism usually affects the whole body without a mosaic distribution (11–13). In the present study, we describe 2 cases of JEB, caused by germline mutations in the LAMB3 gene, which appeared to be revertant mosaics. The skin of the left lower leg of patient 078-01 had initially been fragile but reverted so that it no longer had a tendency to blister due to a higher production of LM-332 trimer. Mutation analysis of distinct biopsies revealed in both patients the presence of at least 5 different somatic second-site LAMB3 mutations that all corrected the same inherited c.628G→A mutation.

Results

Patient 078-01. The proband was a 46-year-old man who consulted our EB center in 1999 with characteristic generalized non-Herlitz JEB features. His clinically normal parents were unrelated, and his siblings were unaffected. There was no family history of inherited skin diseases. Since birth the patient displayed generalized trauma-induced skin blistering that healed with some atrophy and hyperpigmentation. The finger- and toenails were small and short. He had sparse scalp hair with male pattern baldness, mild alopecia of beard hair, and complete loss of eyebrows, eyelashes, and secondary hair. The patient’s teeth all had enamel defects (amelogenesis imperfecta) and had been extracted at the age of 20. Remarkably, the proband claimed that after an erosive period of 7 years, the skin of his left lower leg, which had always shown blisters after minor trauma, healed to clinically unaffected skin that was no longer subject to trauma-induced blistering (Figure 1). The reversion of the phenotype occurred after birth and would therefore be classified as...
The proband’s genomic DNA: the common nonsense mutation c.1903C→T;p.R635X in exon 14; and c.628G→A;p.E210K in exon 7. Since the latter mutation occurs at position –1 of the 5’ donor splice site of intron 7, an effect on mRNA splicing can be expected (14).

To investigate the reversion mechanisms, mutant (reduced expression) and revertant (normal expression) keratinocytes were separately isolated after K140 immunostaining by laser dissection microscopy (LDM). DNA analysis was carried out on the mutation-bearing exons 7 and 14 and on the neighboring exons 6, 8, 13, and 15 (Figure 4A). In the lower leg biopsy IV (R), both mutations, in exon 7 and 14, were still present. Approximate to the germline mutation, we found a second-site mutation in intron 7, c.628+42G→A, 42 nt downstream of the exon/intron 7 border (Figure 4C). Since biopsy III (R) with a mosaic reversion pattern was also taken from the lower leg, we expected to find the same second-site mutation in the revertant keratinocytes. Surprisingly, c.628+42G→A was not present; instead, another nucleotide change — c.596G→C;p.G199A — in exon 7 was observed (Figure 4B). Cloning and sequencing revealed that this additional transversion (c.596G→C) was located on the same allele as the inherited splice site mutation in exon 7. No other nucleotide changes were identified in either of the 2 revertant biopsy specimens. Furthermore, in fibroblasts taken from the same biopsies and in DNA of more than 80 control subjects, the compensatory second-site mutations were absent. Unfortunately, the amount of 25 revertant cells in biopsy II (M) was too low to allow successful DNA analysis. Only 1 sample could be isolated, while we typically perform our sequence analyses on at least 3 separately isolated samples.

To determine the effect of the germline mutation and the additional substitutions on mRNA splicing, we isolated RNA directly from skin sections. cDNA was then synthesized and analyzed by nested RT-PCR using exonic primers that amplified nt 580–848 of LM-332–specific cDNA. In addition to the full-length mRNA transcript (269-bp amplimer), the RNA isolated from affected skin revealed the presence of a small stretch of approximately 25 revertant basal cells that displayed normal LM-332 staining comparable with that of normal age-matched control skin (Figure 2C). Interestingly, both biopsies of unaffected skin showed normal LM-332 staining. Biopsy III (reverted skin [R]) — displayed a mosaic pattern of stretches of normal (3+) and reduced (1+) staining (Figure 2D), while biopsy IV (R) had completely normal 3+ staining (data not shown). Consistent with these findings, 19-DEJ-1 staining was intermittently positive in biopsy III (R) and completely normal in biopsy IV (R).

Electron microscopy of lesional skin demonstrated a subepidermal blister consistent with JEB at the level of the lamina lucida. The blister floor was covered by the lamina densa, and remnants of tonofilaments were present in the blister cavity. Nonlesional affected skin showed a reduced number of hypoplastic hemidesmosomes and less projection of tonofilaments than normal skin (Figure 3A). The lamina densa displayed many duplications and blind “offshoots.” In contrast, the hemidesmosomes were normal in shape in revertant skin (Figure 3B), and the lamina densa showed fewer duplications and offshoots.

Mutation detection was performed on LAMA3, LAMB3, and LAMC2, encoding the 3 chains of LM-332. Sequence analysis revealed compound heterozygosity for 2 mutations in the LAMB3 gene in the proband’s genomic DNA: the common nonsense mutation c.1903C→T;p.R635X in exon 14; and c.628G→A;p.E210K in exon 7. Since the latter mutation occurs at position –1 of the 5’ donor splice site of intron 7, an effect on mRNA splicing can be expected (14).

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![Figure 1](http://www.jci.org)  
**Figure 1**  
Revertant unaffected skin on the left lower leg of proband 078-01 in January 1999. The skin of the left lower leg reverted to clinically healthy skin after an erosive period of 7 years (A, medial aspect; B, lateral aspect), while the skin of the right lower leg remained affected. The area of revertant skin is outlined in black.

late-onset revertant mosaicism (3). On the other leg (right) he developed multiple spinoceullular carcinomas, for which he was treated by amputation in 1998. One year later, metastasis to lymph nodes and lungs occurred, resulting in death of the patient.

Immunofluorescence (IF) antigen mapping of lesional skin biopsy I (mutant skin [M]) showed subepidermal blister formation with the cleavage plane low in the lamina lucida, characteristic of a LM-332–deficient form of JEB. Type XVII collagen was exclusively present in the blister roof and type VII collagen in the blister base. LM-332 was found, although at a reduced level, in both roof and floor. Nonlesional affected skin biopsy II (M) showed reduced binding of LM-332–specific mAbs GB3 and K140 (Figure 2, A and B) and absent binding of 19-DEJ-1 — the marker for JEB — along the whole dermo-epidermal junction. Surprisingly, sections of biopsy II (M) revealed a small stretch of approximately 25 revertant basal cells that displayed normal LM-332 staining comparable with that of normal age-matched control skin (Figure 2C). Interestingly, both biopsies of unaffected skin showed normal LM-332 staining. Biopsy III (reverted skin [R]) — displayed a mosaic pattern of stretches of normal (3+) and reduced (1+) staining (Figure 2D), while biopsy IV (R) had completely normal 3+ staining (data not shown). Consistent with these findings, 19-DEJ-1 staining was intermittently positive in biopsy III (R) and completely normal in biopsy IV (R).

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of 3 aberrant mRNA transcripts: (i) an out-of-frame transcript lacking exon 7; (ii) a transcript with the deletion of exon 7 together with the first 2 nt (AG) of exon 8; and (iii) an in-frame mRNA transcript lacking exon 7 and the first 101 nt of exon 8 (Figure 5, A and B). These 3 alternative mRNA transcripts with deletions of, respectively, 64, 66, and 165 bp, were previously described for the 628G→A mutation by Pulkkinen et al. (14). Detection of their fourth transcript lacking 210 bp was not feasible with our primer set.

Interestingly, a slower migrating amplimer was visible in the RNA isolated from biopsy IV (R). Subsequent cloning and sequencing showed retention of the first 66 bp of intron 7 in this 335-bp product (Figure 5B). In previous work, RNA isolation of mosaic skin biopsy specimens was carried out by staining every fourth tissue section and subsequently using this staining pattern for selection of desired cells from the intermittent sections by LDM isolation (10). Such an approach requires stretches of mutant and revertant cells of sufficient length. Since in biopsy III (R), the areas of reduced and bright staining alternated over short distances — sometimes fewer than 10 basal cells — (Figure 2D), we instead chose to use the whole biopsy for RNA analysis, bearing in mind that about 50% of the cells had normal and 50% reduced LM-332 staining. As expected, RNA of biopsy III (R) had a distribution of mRNA transcripts (Figure 5A, lane 4) different from that of RNA from the affected skin biopsies. The

Figure 3
Normal hemidesmosomes in revertant skin. (A) Ultrastructural examination of the affected skin of proband 078-01 showed a reduced number of abnormal hemidesmosomes. The intermediate filaments were not connected to the flattened basilar cell periphery. (B) The revertant skin of the lower leg reveals intermediate filaments that connect to normal hemidesmosomes in the basilar cell periphery. Black arrows indicate hemidesmosomes and white arrows intermediate filaments. Scale bars: 500 nm.

Figure 4
Identification of the different correcting LAMB3 mutations in patient 078-01. (A) The G→A nucleotide change at position –1 of the 5′ splice site of intron 7 was present in keratinocytes with reduced staining of LM-332. (B) The second-site mutation c.596G→C was present in revertant keratinocytes of biopsy III (R). (C) An additional mutation in intron 7, c.628+42G→A, in revertant keratinocytes of biopsy IV (R). The cryptic splice site, CAG|GT, which is used when the c.628+42G→A substitution is present, is indicated by the dashed line. Red arrows indicate the inherited mutation, and green arrows the second-site mutations. Corresponding amino acid sequences are indicated above the nucleotide sequences.
amplimers that resulted from alternative splicing due to the transi-
tion in the last nucleotide of exon 7 were less abundant (Figure 5B,
transcripts c–e), while the 269-bp amplimer from normal splicing
dominated. Cloning of these normal-sized amplimers showed that
almost all clones carried both c.596G→C and c.628G→A.

Nested RT-PCR was also performed with oligonucleotides ampli-
ifying bp 1,641–2,229 of \textit{LAMB3} cDNA surrounding the other inher-
ited mutation, c.1903C→T; p.R635X. Normal human controls dem-
strated a clear 589-bp PCR product, while in all patient biopsies,
expression of this product was reduced (Figure 5D). An additional
faster-migrating 210-bp amplimer originated from an out-of-frame
transcript with a deletion of exon 14 (379 bp) and is therefore prone
to nonsense-mediated RNA decay. There were no differences in
mRNA or DNA between the unaffected and affected skin biopsies.
Thus, reversion of the inherited p.R635X mutation did not occur.

\textbf{Patient 029-01.} The germline mutation of the second non-Her-
litz JEB proband, 64 years old at the time of publication, has been
described elsewhere (14, 15). His consanguineous parents were
unaffected, but his sister has the same disease. The grandparents
were first cousins. IF of nonlesional affected skin specimens showed
severely reduced LM-332 expression, while type XVII collagen stain-
ing was normal (15). Consistent with this finding, electron micro-
scopic analysis of nonlesional affected skin showed hypoplastic
hemidesmosomes (15). Mutation screening identified the homo-
zygous c.628G→A transition (14).

Because we have observed multiple reversion events in EB patients
(10), we ask patients in our EB clinic on a routine basis whether they
have patches of clinically unaffected skin. In early 2006, patient 029-01
indicated several unaffected regions at his arms, shoulder, and chest
(Figure 6). Unfortunately, he could not remember how long these
patches had been present and whether they had increased in size
over time. Older photographic material was lacking. IF microsco-
py of all 4 biopsies of unaffected skin showed keratinocytes with
bright staining comparable to that of normal control skin (data not
shown), thereby confirming that this EB patient was another indi-
vidual with revertant mosaicism. Consistent with this finding, in
the revertant skin, the number of hemidesmosomes was normal, in
contrast to the reduced amount in the mutant skin.

Subsequent DNA analysis of the revertant keratinocytes by nest-
ed PCRs of exons 5–9 revealed different second-site mutations in
The inherited G → A mutation generates 4 additional aberrant transcripts as described by Pulkkinen et al. (14). RNA splicing is directed by the 5′ donor splice site, the 3′ acceptor splice site, and the branch point sequence that is located 18–40 nt upstream of the acceptor splice site (reviewed in ref. 21). In the human genome, these splicing sequences are poorly conserved; only the GT at the 5′ end of the intron, AG at the 3′ end of the intron, and branch point adenosine at the 2′ position are almost invariable. Since natural splice sites can be different from the consensus sequence, both splice site strength and accessory regulatory sequences influence splice site selection. Various web-based resources for splice site prediction are available, such as Automated Splice Site Analyses, located at https://splice.cmh.edu. Computational analyses of our LAMB3 genomic sequences showed that the c.628G → A mutation weakens the individual information content (R) of the natural donor site from 8.2 to 5.1 bits. Besides this 5′ splice site, 2 other donor sites with a high R value were present in the analyzed sequence. These sites were TGG|GT, with an R of 4.3 bits, and CAG|GT, with an R of 3.2 bits, located 37 and 66 nt downstream of the natural exon/intron 7 border, respectively.

The second-site mutation c.628+42G → A lowered the R value of the alternative donor site TGG|GT drastically from 4.3 to 0.8 bits, whereas the CAG|GT site remained unchanged at 3.2 bits. The weakening of the TGG|GT donor site might favor use of the cryptic CAG|GT site by the spliceosome, thereby resulting in the observed larger mRNA transcript with retention of the first 66 nt of intron 7. The in-frame insertion led to incorporation of a stretch of 22 amino acids — SQCGYFSCPWNYGRKRKNWSP — in the N-terminal domain of the β3 chain. Although this stretch contained 2 amino acids with basic side chains, 4 with acidic side chains, and 6 with bulky aromatic side chains, the resulting LM-332 protein was apparently functional, as it resulted in reversion of the skin phenotype.

The second-site mutation c.596G → C changed a glycine to alanine at amino acid position 199, and second-site c.619A → C changed a lysine to glutamine at position 207. More importantly, both also affected mRNA splicing, as more normal-sized transcripts were present in the revertant cells. Computational analyses did not demonstrate effects on R values of the splice sites used. We also excluded the possibility that the mutations altered exonic splicing enhancer (ESE) sequences using RESCUE-ESE software (http://genes.mit.edu/burgelab/ rescue-ese). ESEs enhance splicing when present downstream of a 3′ splice site and/or upstream of the 5′ splice site (22). A third factor that may influence splice site selection is a possible effect of these second-site mutations on the RNA secondary structure that contains the c.628G → A mutation. Single nucleotide alterations can affect the secondary structure of RNA, which in turn can influence RNA splicing (23, 24).

Two nucleotides downstream of the wild-type 3′ acceptor site of intron 7, CAG|AG, a cryptic splice site, GAG|GT, was present. Therefore c.628G → A generated not only a 64 bp–deleted transcript, but also the 66 bp–deleted transcript. The second-site mutation 629-1G → A reduced the strength of the natural acceptor site from 10.5 to 2.9 bits, whereas the cryptic splice site increased in strength from 6.0 to 7.8 bits. RT-PCR analysis indeed demonstrated preferential use of the cryptic splice site generating an in-frame rather than an out-of-frame transcript. Translation resulted in a smaller functional β3 polypeptide lacking 22 amino aci...
acids within the N-terminal domain of the short arm. The final second-site mutation, c.565-3T→C, was located in the 3′ acceptor site of intron 6 and gave a small increase, from 5.3 to 5.6 bits, in the $R_i$ value. Although small, it might explain the higher production of wild-type mRNA transcript.

The other inherited mutation, p.R635X, which is the most common mutation in JEB patients of European origin, predicts a premature termination codon (PTC) within the coiled-coil rod of the LM-332 protein. The corresponding transcript level is expected to be markedly reduced due to nonsense-mediated RNA decay. RT-PCR around exon 14 indeed showed reduced expression of the 589 amplimers as described by Pulkkinen et al. (25). Also, a smaller migrating 210-bp amplimer was detected for patient 078-01 belonging to an out-of-frame transcript lacking exon 14 (379 nt). Although this nonbeneficial transcript has not been reported before, it is not uncommon that PTCs induce exon skipping (26).

Figure 7
The inherited homozygous c.628G→A mutation was present in 029-01 proband’s keratinocytes with normal (A–C) as well as reduced (D) LM-332 staining. (A) In biopsy I (R), a second-site mutation, c.619A→C;p.K207Q, was present in exon 7. Biopsy III (R) had an additional substitution, c.565-3T→C, in the 3′ splice site of intron 6 (B) and biopsy IV (R) an additional c.629-1G→A change in the 3′ splice site of intron 7 (C). (D) None of these additional substitutions were seen in LDM-isolated mutant keratinocytes. Red arrows indicate the inherited mutation, and green arrows the second-site mutations. Amino acid sequences are indicated above the nt sequences.
Arguments in favor of random mutagenesis. The average mutation rate in humans is estimated to be 175 mutations per diploid genome per generation by Nachman and Crowell (31). It may well be that our patients have a higher mutation rate and that therefore different advantageous second-site mutations accumulated in keratinocytes. Other mutations in different genes or in different cell types that are not advantageous for the cell may just get lost. Accordingly, no second-site mutations were detected in peripheral blood or the fibroblast samples. Such an increased mutation rate can be the result of inactivation of a caretaker gene (32). Inactivation results in genetic instabilities, causing an increased mutation rate affecting all genes. In light of this hypothesis, it is interesting to note that both patients developed cancer, which is known to result from an accumulation of somatic mutations (33).

Arguments in favor of a directed mutagenesis. In the heterozygous patient 078-01, all 3 second-site mutations correct the same 628G→A mutation. No second-site mutation was observed for the primary 1903C→T mutation on the other allele. The FAH gene patient described by Bliksrud et al. (16) was also heterozygous, and similar to our findings, both in vivo reversions were located on the same allele. No knowledge of the proper wild-type sequence is required to execute the reversion repair mechanism. While it can be argued that in a heterozygous patient, information on the correct sequence is available on the other allele, this cannot be the case in a homozygous patient. In 029-01, homozygous for 628G→A, 4 different second-site mutations were identified. Also, the probands described by Wada et al. (17) with 6 different reversions and by Rieux-Laucat et al. (27) with 3 different reversions were both homozygous. More cases of in vivo reversion in homozygous and hemizygous patients have been described previously (2).

Recently, we reported revertant mosaicism for type XVII collagen non-Herlitz JEB (10). In these COL17A1 mosaic patients, the revertant patches remained stable during life and did not expand. Apparently, the revertant stem cells did not have a selection advantage compared with their deficient counterparts. Therefore, we concluded that the correcting mutations leading to the healthy skin patches of tens of square centimeters in size occurred during embryogenesis. In the LAMB3 mosaic patients described here, the situation was different. According to patient 078-01, his revertant skin patch increased in size, while patient 029-01 had not noted extension of the healthy skin area. This difference might be explained by the fact that patient 029-01 (c.628G→A/c.628G→A) likely has a higher level of LM-332 production in his deficient cells than patient 078-01 (c.628G→A/p.R635X), because the allele containing the nonsense mutation in the latter was not contributing to the LM-332 production. Therefore, expansion of reverted keratinocytes could indeed have been easier in proband 078-01 than in 029-01, as the deficient cells were less able to compete. LM-332 is involved in cell locomotion and migration in wound healing (34, 35). Interactions with α2β1 and α3β1 are important for cell attachment, spreading, and migration, whereas binding to α6β4 results in stable anchorage without cell spreading (36, 37). Also, suppression of endogenous LM-332 in an oral squamous cell carcinoma cell line led to decreased cell attachment and increased migration (38). The benefit of reversion is possibly related to the role of LM-332 in migration; however, this potential selection advantage of LM-332 revertant stem cells requires thorough investigation.

The treatment for genetic diseases seems to lie in gene therapy. Mavilio et al. (5) recently showed in a phase I clinical trial the correction of LM-332–deficient JEB by transduction of retroviral
vector expressing β3 cDNA. A 36-year-old man received a transplant of cultured epidermal sheets on both legs after removal of the outer skin layer. Transplantation was successful, as during the first year of follow-up, blistering, infections, inflammations, and an immune response were absent. Revertant mosaicism opens the fascinating possibility of “revertant cell therapy” for mosaic patients using patient’s own naturally corrected cells for transplantation. In LAMB3 revertant mosaic patients, one might take advantage of the patient’s own naturally corrected cells for skin transplantation. This autologous cell therapy bypasses the phase of molecular gene correction. Revertant mosaicism was thought to be a rare event, but our recent observations indicate that it might occur at a higher frequency than expected.

**Methods**

**Biopsy sites.** From patient 078-01 four 4-mm punch biopsies were taken and snap frozen for IF microscopy: 1 from lesional affected skin from the left upper arm (biopsy I [M]), 1 from nonlesional affected skin from the left upper arm (biopsy II [M]), and 2 from unaffected skin of the left lower leg (biopsies III [R] and IV [R]). For electron microscopy, three 2-mm punch biopsies were taken: 1 from lesional affected skin of the left upper arm, 1 from nonlesional affected skin of the left upper arm, and 1 from unaffected skin of the left lower leg.

The second proband, 029-01, has been described elsewhere, specifically, as patient no. 9 in the publication of Jonkman et al. (15) and as patient no. 2 in the publication of Pulkkinen et al. (14). Six 4-mm punch biopsies were taken for IF microscopy: 1 from unaffected skin of the right shoulder (biopsy I [R]), 1 from unaffected skin of the right upper arm (biopsy III [R]), 1 from unaffected skin of the right lower arm (biopsy III [R]), 1 from nonlesional affected skin of the left upper arm (biopsies V [M] and VI [M]). For electron microscopy, two 2-mm punch biopsies were taken: 1 from nonlesional affected skin of the left upper arm and 1 from unaffected skin of the right upper arm. Informed consent for the scientific use of the material and photographs was obtained from both patients according to the guidelines of the ethics committee of the University Medical Center Groningen and the Declaration of Helsinki.

**Immunomorphological analysis of skin and cultured keratinocytes.** Details of IF microscopy, electron microscopy, and keratinocyte culturing have been extensively documented elsewhere (39). For detection of LM-332, two antibodies were used: K140 specific to the β3 chain (gift from B. Burgeson, Harvard University, Boston, Massachusetts, USA) and GB3, recognizing a conformational epitope of the γ2 chain (Abcam). As a marker for JEB, we used mAb 19-DEJ-1 (gift from J.D. Fine, Vanderbilt University School of Medicine, Nashville, Tennessee, USA). Type XVII collagen was stained with 1A8C and 1D1 (gifts from K. Owaribe, Nagoya University, Nagoya, Japan), type VII collagen with LH7:2 (gift from I. Leigh, St. Bartholomew’s and the Royal London School of Medicine and Dentistry, London, United Kingdom). The Alexa Fluor 488–conjugated goat anti-mouse IgG antibody (Molecular Probes) was used as secondary step with labeled conjugate.

**DNA and RNA isolation from skin sections.** DNA recovery with LDM was performed as described previously (10). Monoclonal antibody K140 was used to differentiate between keratinocytes with normal and reduced staining. For DNA isolation, about 200 cells were collected in a 0.2-ml reaction tube. Proteinase K digestion was for 60 minutes at 55°C, followed by proteinase K inactivation at 98°C for 15 minutes. The final aliquots were used for

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**Figure 9**

Schematic drawing showing the different second-site mutations that all corrected LAMB3:c.628G→A. (A) The inherited germline mutation c.628G→A is depicted as black rectangles, while the second-site mutations are depicted as white rectangles. Cells with a mutant phenotype are white, and those with a revertant phenotype green. (B) Distribution of the mutations in the LAMB3 gene. The red arrow indicates the inherited mutation, and green arrows indicate the second-site mutations.
PCR. For RNA isolation, three 10-μm skin sections were lysed in 100 μl of lysis buffer plus 0.7 μl β-mercaptoethanol. Total RNA (15 μl) was then prepared using the Stratagene RNA Microprep kit. cDNA was synthesized as described elsewhere (10).

**Identification of the mutations in LDM samples.** For LAMB3 mutation detection in LDM-isolated DNA, we used nested PCR. One microliter of the first PCR was used for the second PCR. PCR cycling conditions were 5 minutes at 94°C, followed by 35 cycles at 94°C for 45 seconds, 55°C for 45 seconds, and 72°C for 1 minute, and a final extension at 72°C for 7 minutes. Water was used as a negative control. Primer sequences were designed to amplify the different exons of LAMB3 and expected product sizes are listed in Supplemental Table 1 (supplemental material available online with this article; doi:10.1172/JCI39465DS1). After PCR, aliquots of 14 μl were examined on 1.5% agarose gels. The Department of Medical Genetics of the University Medical Center Groningen provided genomic DNA of 85 control subjects to exclude the possibility that detected second-site mutations were rare polymorphisms. All donors gave informed consent. Also, CDNA samples from dissected cells were subjected to nested PCR. All primers were designed in such way that the PCR product contained sequences from multiple exons (Supplemental Table 2). All PCR analyses were repeated with templates from at least 3 separate LDM-obtained nucleic-acid isolations. The resulting PCR products were cloned into the pCR4-TOPO vector (Invitrogen). For each revertant biopsy specimen, more than 10 clones were selected and sequenced.

**Accession numbers.** The GenBank accession numbers for the human LAMB3 gene are U17744–U17760.

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