



Molecular Therapy

Original Article

CRISPR/Cas9-Mediated *In Situ* Correction of LAMB3 Gene in Keratinocytes Derived from a Junctional Epidermolysis Bullosa Patient

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Deficiency of basement membrane heterotrimeric laminin 332 component, coded by *LAMA3*, *LAMB3*, and *LAMC2* genes, causes junctional epidermolysis bullosa (JEB), a severe skin adhesion defect. Herein, we report the first application of CRISPR/Cas9-mediated homology direct repair (HDR) to *in situ* restore LAMB3 expression in JEB keratinocytes *in vitro* and in immunodeficient mice transplanted with genetically corrected skin equivalents. We packaged an adenovector carrying Cas9/guide RNA (gRNA) tailored to the intron 2 of *LAMB3* gene and an integration defective lentiviral vector bearing a promoterless quasi-complete *LAMB3* cDNA downstream a splice acceptor site and flanked by homology arms. Upon genuine HDR, we exploited the *in vitro* adhesion advantage of laminin 332 production to positively select LAMB3-expressing keratinocytes. HDR and restored laminin 332 expression were evaluated at single-cell level. Notably, monoallelic-targeted integration of *LAMB3* cDNA was sufficient to *in vitro* recapitulate the adhesive property, the colony formation typical of normal keratinocytes, as well as their cell growth. Grafting of genetically corrected skin equivalents onto immunodeficient mice showed a completely restored dermal-epidermal junction. This study provides evidence for efficient CRISPR/Cas9-mediated *in situ* restoration of LAMB3 expression, paving the way for *ex vivo* clinical application of this strategy to laminin 332 deficiency.

INTRODUCTION

Junctional epidermolysis bullosa (JEB) is a recessively inherited skin disorder characterized by fragility of the skin and mucous membranes and by tissue separation within the lamina lucida of the cutaneous basement membrane zone (BMZ). The majority of JEB cases are caused by defects in laminin 332, the major adhesion ligand of basal epithelial cells expressed in the BMZ of the skin and mucous epithelia. Laminin 332 is a multi-domain glycoprotein formed by 3 subunits, $\alpha 3$, $\beta 3$, and $\gamma 2$, coded by *LAMA3*, *LAMB3*, and *LAMC2* genes, respectively. Restored expression of laminin 332 protein in defective skin keratinocytes by gene addition strategies results in a complete re-establishment

of the adhesion property of keratinocytes *in vitro*^{1,2} and in a permanent correction of a normal dermal-epidermal junction *in vivo*.^{3,4}

The gene addition strategy based on viral (e.g., adeno-associated virus [AAV] and lentiviral and retroviral vectors) delivery of an expression cassette for the correct gene has been the strategy of choice to treat severe genetic disorders, e.g., junctional^{3,4} and dystrophic epidermolysis bullosa (DEB),⁵ hemophilia B,⁶ and X-linked severe combined immunodeficiency (SCID-X1).⁷ The gene addition strategy has shown excellent results over the years in many cell contexts and with multiple vectors and genes. However, permanent correction of disease-causing mutations or locus-specific insertion of a therapeutic cDNA remains preferable to allow long-term physiological and tissue-restricted expression of the introduced sequence.

In the last years, emerging genome-editing strategies, able to modify nucleic acids within disease-affected cells and tissues, have shown therapeutic potential for the treatment of monogenic, highly penetrant diseases, such as SCID-X1⁸ and hemophilia B.⁹ Among the four major classes of nucleases, meganucleases,¹⁰ zinc finger nucleases (ZFNs),¹¹ transcription activator-like effector nucleases (TALENs),¹² and CRISPR-associated nuclease Cas9,¹³ CRISPR/Cas9 system from *S. pyogenes* (SpCas9) clearly boosted the applications of genome editing to gene therapy. It has turned out that Cas9-mediated homologous recombination (HR) of a donor DNA template offers a valuable option to correct the mutated genes associated to monogenic disorders.^{14,15} This study aims for the first time to apply CRISPR/Cas9-triggered HR to *in situ* re-establish LAMB3 expression by integrating an almost complete *LAMB3* cDNA. We employed, as therapeutic donor, a promoterless *LAMB3* cDNA engineered to be transcriptionally silent

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unless a splicing event between genomic exon 2 and donor exon 3 occurs, upon HR event. This donor design overcomes most of the safety concerns of gene therapy and holds several advantages: (1) site-specific integration with transgene regulation and transcription controlled by the endogenous promoter; (2) quasi-complete *LAMB3* cDNA (exon 3–23), thus tackling most of the mutations mapped in JEB patients; (3) lack of a promoter in the donor template vector, thus reducing the probability of activating neighboring oncogenes if undesired donor integration occurred; and (4) canonical splice acceptor site of *LAMB3* gene conferring high expression of laminin 332, which is required to restore the *in vitro* adhesion property of corrected keratinocytes. Restoration of laminin 332 expression in the keratinocyte culture has been demonstrated to provide a biological system to positively select gene-corrected cells,¹⁶ avoiding cell sorting that could impair cell viability, or antibiotic selection, not feasible in many clinical applications. Thus, exploiting the *in vitro* adhesion advantage of laminin-332-corrected keratinocytes, we propose a novel and safe gene editing strategy to *in situ* restore *LAMB3* expression upon canonical splicing event in the defective gene of JEB keratinocytes.

In this study, a *LAMB3* editing strategy was applied to keratinocytes from two JEB patients: the first carrying a homozygous frameshift mutation (c.1945dupG) in the exon 14 of *LAMB3* gene, leading to the formation of a premature stop codon within exon 15 (p.A649fsX38),¹⁷ and the second carrying a frequent (45%–63%) homozygous c.1903C > T transition in the exon 14 of *LAMB3* gene, resulting in an Arg635-to-ter (R635X) substitution.¹⁸ To restore the expression of wild-type *LAMB3* gene, we designed two molecular tools: a guide RNA (gRNA) for SpCas9 targeting *LAMB3* intron 2 and a promoterless *LAMB3* cDNA (exon 3–23) flanked by a splice acceptor and a poly-adenylation (pA) sites and embedded between two homology arms (HAs) flanking the target site. JEB keratinocytes were co-transduced with an adenoviral vector carrying expression cassettes for SpCas9 and gRNA (AdCas9gRNA) and non-integrating lentiviral vector bearing the therapeutic donor template (IDLV-L.B3). Transduced cells were positively selected *in vitro* exploiting physiological adhesion advantages of gene-corrected cells. Molecular analysis on single isolated clones demonstrated an *in situ* monoallelic and biallelic integration of the therapeutic donor template restoring the expression of laminin 332 heterotrimer. *LAMB3*-corrected clones showed morphology and adhesive properties comparable to wild-type (WT) keratinocytes and increased proliferation with respect to JEB cells. A functional reconstruction of dermal-epidermal junction was demonstrated in immunodeficient mice transplanted with skin equivalents from corrected clones. This study represents the first proof that *in situ* integration of wild-type *LAMB3* cDNA spanning from exon 3 to 23 is able to correct the JEB phenotype *in vitro* and in a mouse model.

RESULTS

Correction of Keratinocytes from a JEB Patient by CRISPR/Cas9-Mediated Homology Direct Repair of Promoterless *LAMB3* cDNA

JEB disease exhibits heterogeneity in the location and number of mutations in *LAMB3* gene. To tackle a wide range of JEB-causative

mutations spread all along the transcription unit of *LAMB3* gene, we employed CRISPR/SpCas9 system. We designed a gRNA tailored to the second intron of *LAMB3* gene to trigger site-specific HR of a promoterless *LAMB3* cDNA (Figure 1A). The therapeutic *LAMB3* donor carried the splice acceptor site of exon 3 followed by a super exon consisting of correct cDNA from exon 3 to 23 and a bovine growth hormone (bGH) pA site. This promoterless therapeutic donor cassette, flanked by HAs, was cloned in reverse orientation into an integration defective lentiviral vector (IDLV-L.B3) in order to avoid cryptic splicing site in 5' long terminal repeat (LTR),¹⁹ leading to expression of truncated proteins (Figure 1A). SpCas9 and gRNA were cloned into a first-generation adenoviral vector (AdCas9gRNA) able to efficiently transduce human keratinocytes.^{20,21}

The nuclease-mediated *in situ* integration of the therapeutic *LAMB3* donor was addressed in immortalized keratinocytes derived from a JEB patient carrying the c.1945dupG *LAMB3* mutation (referred to as JEB.A649fsX38). This homozygous mutation maps on exon 14 and results in a frameshift starting at codon 649 and leading to the formation of a premature stop codon within exon 15 (p.A649fsX38).¹⁷ As a consequence, the mutant mRNA is degraded by nonsense-mediated mRNA decay mechanism and, in case of occurring protein synthesis, translated in a β chain truncated in the rod domain, which is required for laminin-332 assembly. Thus, the mutation leads to a complete loss of *LAMB3* expression.

We first transduced JEB.A649fsX38 keratinocytes with increasing doses of AdCas9gRNA vector (MOI 10^4 , 2×10^4 , 5×10^4 , and 10^5 vector genomes [vg]/cell) to evaluate gRNA editing efficiency. Surveyor assay of AdCas9gRNA-transduced cells clearly showed editing at all tested MOIs (Figure S1). To determine indels frequency, tracking of indels by decomposition (TIDE) analysis²² on PCR amplicons derived from transduced JEB.A649fsX38 cells was performed and showed 25%, 41%, 67%, and 76% ($p < 0.05$) of indels, respectively (Figure S2). Indels types upon AdCas9gRNA delivery (MOI 2×10^4 vg/cell) was more accurately evaluated by bioinformatics analysis (D.B., V. Marigo, and A.R., unpublished data) of single Sanger sequences (Figure S3) and resulted in 56.3% total indels, in particular 40.8% insertion, 14.1% deletion, and 1.4% mismatches. To note, higher doses resulted in toxic effect; thus, the MOI 10^4 and 2×10^4 vg/cell were selected for *LAMB3*-editing experiment. We co-transduced JEB.A649fsX38 keratinocytes with AdCas9gRNA (MOI 10^4 and 2×10^4 vg/cell) and IDLV-L.B3 (4.32×10^2 vg/cell) vectors. Targeted integration (TI) PCR confirmed that homology direct repair (HDR) events precisely occurred, as demonstrated by attended bands at both 5' and 3' vector-genome junctions (Figure 1B). However, flow cytometric analysis of laminin 332 expression in co-transduced JEB.A649fsX38 keratinocytes showed that, as previously observed,²⁰ the HDR frequency was barely detectable (% *LAMB3*⁺ cells = 0.48 ± 0.19 ; Figures 1C and S4), remarking the requirement of positively selecting *LAMB3*-expressing cells.

To predict the most likely off-target sites of gRNA, we used public web servers (<http://crispr.mit.edu>; <https://crispr.dbcls.jp>)²³ to assess

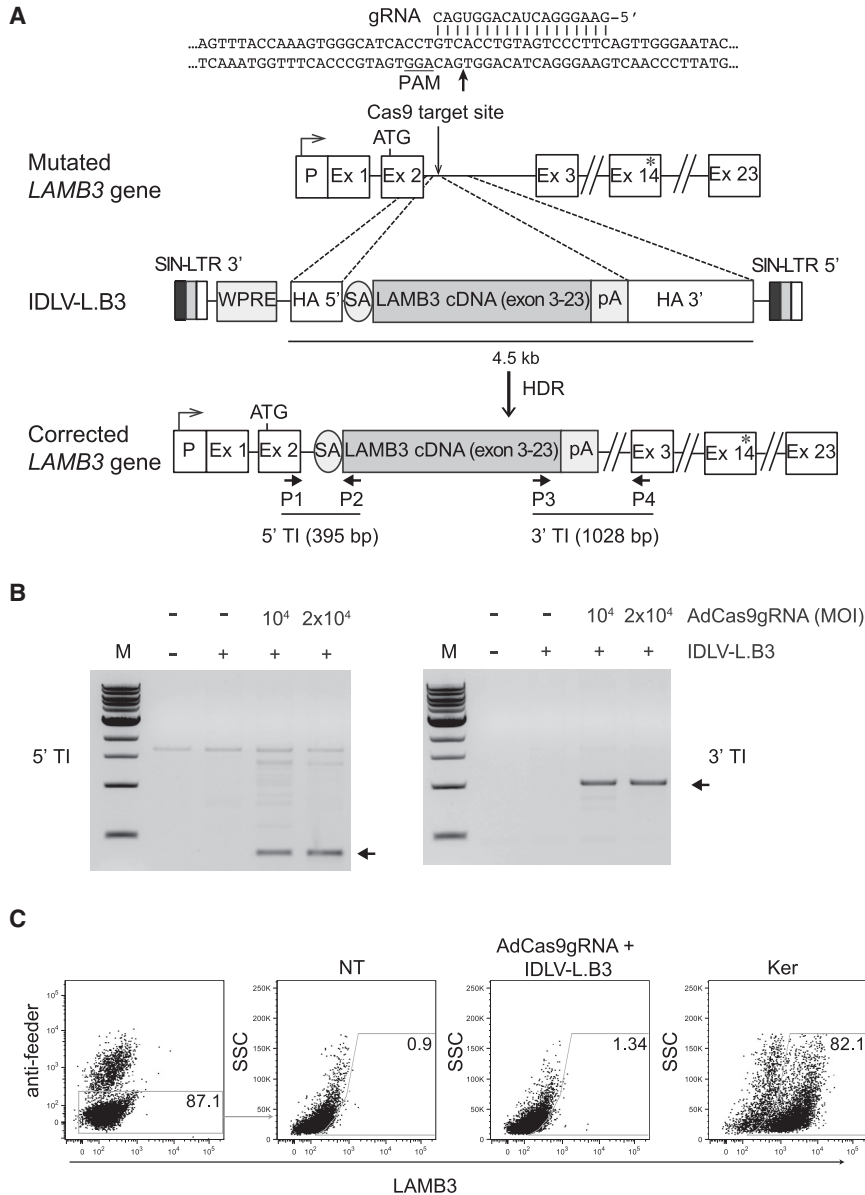


Figure 1. CRISPR/Cas9-Mediated HDR into Intron 2 of LAMB3 Gene in Keratinocytes Derived from a JEB Patient

(A) (Top) Endogenous mutated *LAMB3* gene carrying 23 exons (Ex) is schematized (not in scale). The arrow above the promoter region (P) indicates the transcription start site; Cas9 target site and the ATG are depicted. Sequences of the selected gRNA and PAM on intron 2 are reported. The asterisk indicates *LAMB3* mutations generating a stop codon. (Middle) Schematic representation (not in scale) of promoterless *LAMB3* donor (4.5 kb in length) cloned in opposite orientation into IDLV backbone, generating IDLV-L.B3 vector, is shown. (Bottom) HDR event into the target site re-establishing the transcription of corrected *LAMB3* cDNA is shown. HA, homology arm (90 bp HA5' and 413 bp HA 3'); pA, polyadenylation site (278 bp); SA, splicing acceptor; SIN-LTR, self-inactivating long terminal repeat (LTR); WPRE, post-transcriptional regulatory element of woodchuck hepatitis virus. P1–4 represent primers used for targeted integration and allele discrimination PCRs. 5' (395 bp) and 3' (1,028 bp) TIs indicate the size of amplicons generated upon genuine HDR event. (B) PCR analyses on genomic DNA from JEB.A649fsX38 cells transduced with IDLV-L.B3 (4.32×10^2 vg/cell) in absence or in presence with AdCas9gRNA at indicated MOIs (vg/cell) or not transduced. Two couple of primers specific for 5' and 3' vector genome junctions (P1+P2 and P3+P4, respectively, as depicted in A) amplified 395- and 1,028-bp bands indicated by arrows, respectively, are shown. M, 1 kb molecular weight marker. (C) Flow cytometric analysis of laminin 332 expression in JEB.A649fsX38 cells not transduced (NT), co-transduced with IDLV-L.B3 (4.32×10^2 vg/cell) + AdCas9gRNA (MOI 10^4 vg/cell), and in WT keratinocytes (Ker), after feeder exclusion (left panel). The boxed gates represent the percentage of *LAMB3*-positive keratinocytes (0.9%, 1.34%, and 82.1%).

Selective Advantage of *LAMB3*-Corrected JEB Keratinocytes *In Vitro*

Taking advantage of *in vitro* adhesion properties of *LAMB3*-expressing cells,¹ we selected bulk populations of *LAMB3*-corrected keratinocytes by 2 rounds of cell detachment from tissue culture plates using a trypsin solution. The restored expression of *LAMB3* gene conferred to JEB.A649fsX38 keratinocytes an increased adhesion to a 3T3 feeder layer seeded onto the tissue culture plate with respect to untreated cells and to JEB.A649fsX38 treated with IDLV-L.B3 alone. Indeed, qRT-PCR analysis before selection showed a barely detectable increase of *LAMB3* mRNA in co-transduced bulks compared to control JEB.A649fsX38. Conversely, after two rounds of selection, *LAMB3* expression in co-transduced JEB.A649fsX38 cells was a 6- or 7-fold higher respect to untreated cells (Figure 2A). We further evaluated the enrichment of *LAMB3*-corrected keratinocytes by flow cytometry of laminin 332 expression. Up to 17% laminin-332-positive cells were detected in co-transduced JEB.A649fsX38

and prioritize potential Cas9 activity at off-target loci based on predicted positional bias of a given mismatch in gRNA protospacer sequence and total number of mismatches to the intended target site. The CRISPR design tool scored a total of 186 potential off-target sites in the human genome (data not shown). The top 7 potential off-target sites (Table S1) were assessed by Surveyor assay and TIDE analysis on JEB.A649fsX38 cells transduced at $MOI 2 \times 10^4$ vg/cell. None of the 7 predicted off-target loci showed detectable levels of off-target gene modification by Surveyor assay (Figure S5A). TIDE analyses scored a 0.9% frequency of indels in the OT7, although no significant ($p < 0.05$) level of indels was measured for the other off targets (OTs) (Figure S5B).

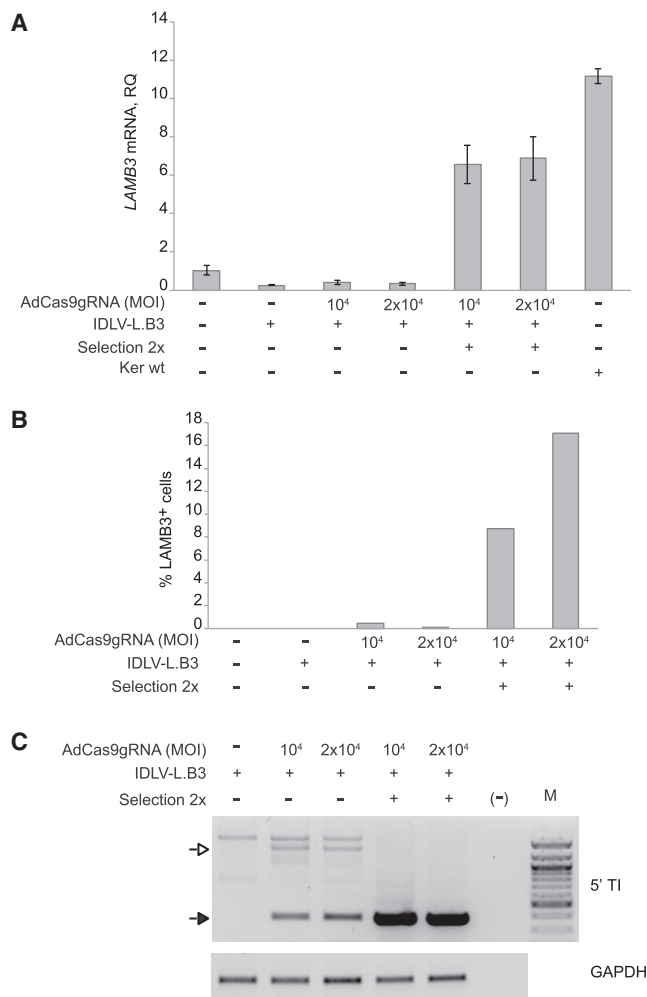


Figure 2. Restored LAMB3 Expression in Gene-Corrected Keratinocytes Selected by *In Vitro* Adhesion Strategy

(A) qRT-PCR analysis on *LAMB3* expression in JEB.A649fsX38 keratinocytes bulks transduced with IDLV-L.B3 (4.32×10^2 vg/cell) alone or in combination with AdCas9gRNA vector at increasing MOIs (vg/cell). *LAMB3* expression was measured in JEB.A649fsX38 cells before and after 2 rounds of selections and in WT keratinocytes, as control. Data are representative of three independent PCR amplifications (mean \pm SEM; n = 3). (B) Representative experiment of flow cytometric analysis of LAMB3⁺ cells before and after selection of JEB.A649fsX38 keratinocytes co-transduced with IDLV-L.B3 (4.32×10^2 vg/cell) and AdCas9gRNA (MOI 10⁴ and 2 \times 10⁴ vg/cell). The frequency (%) of LAMB3-positive cells was determined after subtraction of background measured in not transduced cells. (C) Genomic analysis of 5' TI in pre- and post-selected JEB.A649fsX38 keratinocytes co-transduced with IDLV-L.B3 (4.32×10^2 vg/cell) and AdCas9gRNA (MOI 10⁴ and 2 \times 10⁴ vg/cell). The size of PCR bands corresponding to genuine HR (395 bp, black arrow) and HR-independent (approximately \sim 1,400 bp, white arrow) integration of the donor template is indicated. The bottom gel shows PCR amplification targeting the *GAPDH* (glyceraldehyde-3-phosphate dehydrogenase) gene. M, 100-bp molecular weight marker.

bulks with respect to untreated JEB.A649fsX38, indicating a robust enrichment of *LAMB3*-corrected cells after 2 rounds of selection (Figure 2B).

The positive selection was also confirmed at genomic level by 5' TI PCR performed pre- and post-2 rounds of detachment (Figure 2C), pointing out that the selective adhesion procedure enriched cells undergone genuine HR event and splicing between the genomic splice donor site in exon 2 and the splice acceptor site of exon 3 cloned upstream *LAMB3* cDNA. As shown in Figure 2C, the adhesion strategy negatively selected the HR-independent integration of the donor template represented by an \sim 1,400-bp PCR band amplified in co-transduced JEB.A649fsX38 cells pre-selection. Sequencing of the PCR band indicated the *in situ* integration of the 3' end of IDLV vector (Figure S6).

The selective advantage of *LAMB3*-corrected cells was also demonstrated in immortalized JEB keratinocytes carrying the R635X mutation (referred to as JEB.R635X),¹⁸ which is responsible for complete loss of laminin 332 expression. JEB.R635X keratinocytes were transduced with AdCas9gRNA (MOI 2×10^4 vg/cell) in presence or absence of IDLV-L.B3 (4.32×10^2 vg/cell). Three days post-transduction, we investigated the presence and frequency of indels in the target site by Surveyor assay and TIDE analysis. The results reported in Figures S7A and S7B showed a 56.7% of indels in JEB.R635X keratinocytes transduced with AdCas9gRNA vector. Co-transduction of JEB.R635X cells resulted in expected HDR event as demonstrated by 5' and 3' TI PCRs (Figure S7C). After 2 rounds of selection, we verified the enrichment of *LAMB3*-corrected cells by 5' TI PCR, qRT-PCR, and flow cytometry as previously performed in JEB.A649fsX38 keratinocytes pre- and post-selection. The results (Figures S7D–S7F) demonstrated the positive selection of cells undergoing genuine HR events and the restored *LAMB3* expression in co-transduced JEB.R635X keratinocytes respect to control cells.

Restored *LAMB3* Expression in Selected Single-Cell Clones

To molecularly characterize the *in situ* integration of *LAMB3* cDNA, enriched JEB.A649fsX38 keratinocytes were cloned by limiting cell dilution. Twenty-four out of 54 clones derived from AdCas9gRNA (MOI 10⁴ vg/cell) infection and 15 out of 34 clones from higher dose (MOI 2×10^4 vg/cell) resulted positive in 5' and 3' TI PCR analyses (Figures S8A and S8B), indicating that more than 2 rounds of selective adhesion would be required to fully enrich for *LAMB3*-corrected clones. Remarkably, the percentage of corrected clones was 44% from both bulk populations, indicating that AdCas9gRNA-tested doses did not influence the frequency of HR. Eight out of 39 clones, equally distributed in the two bulk keratinocytes, showed a 3' TI PCR longer than expected (Figures S9A and S9B). Deeper investigation of this aberrant HR event revealed an *in situ* integration of the 5' end of the IDLV vector (Figures S9C and S9D). Nevertheless, these clones correctly expressed *LAMB3* mRNA (Figure S10A). For further analysis, we expanded 28 *LAMB3*-corrected clones and 10 negative clones for targeted integration PCR assay. First, we analyzed allelic distribution of targeted clones with a PCR-based allelic discrimination strategy employing 3 primers (P1, P2, and P4) schematized in Figure 1A. Twenty-four and four targeted clones displayed monoallelic and biallelic integration of the therapeutic donor, respectively (Figures 3A and S10B), meaning that, in the vast majority of clones, only one allele

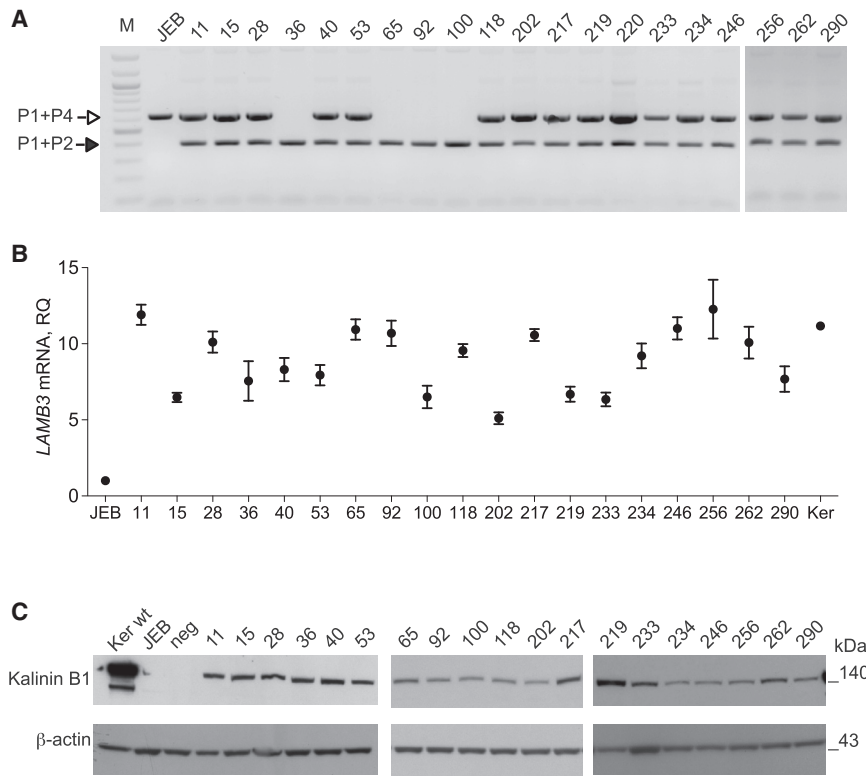


Figure 3. Molecular Characterization of LAMB3-Corrected Clones

(A) Allelic discrimination: a three-primer-based (P1, P2, and P4; see Figure 1A) PCR approach was employed to discriminate between parental allele (628 bp; P1+P4; white arrow) and targeted allele (395 bp; P1+P2; black arrow). Monoallelic clones hosting one copy of the therapeutic donor showed both PCR bands. Biallelic clones hosting two copies of the therapeutic donor displayed only 395-bp band. M, 100-bp molecular weight marker. (B) qRT-PCR analysis on *LAMB3* expression in representative clones with respect to untreated JEB.A649fsX38 keratinocytes and WT keratinocytes (Ker). Data are representative of three independent PCR amplifications (mean \pm SEM; n = 3). (C) Western blot analysis showing laminin 332 expression in lysates from representative clones using an antibody against Kalinin B1 (an alternative name of laminin subunit beta-3 protein). β -actin staining served as loading control. Wild-type keratinocytes (Ker wt) and JEB.A649fsX38 keratinocytes (JEB) are used as positive and negative controls, respectively. Neg. indicates a negative clone. Protein molecular weights are indicated (kDa).

carrying the homozygous mutation hosted the therapeutic donor. Laminin 332 expression was rescued upon splicing between genomic and donor template splice sites in all 5' and 3' TI-positive clones as indicated by qRT-PCR (Figure 3B) and western blot analysis (Figure 3C) reported for representative 19 clones. Laminin 332 expression, both in terms of RNA and protein level, was variable but always stronger than in negative clones or JEB.A649fsX38 cells, ranging from 5- to 12-fold (Figure 3B). Microscopic appearance of representative corrected and control clones seeded on a feeder layer confirmed that *LAMB3*-corrected clones formed colonies comparable to normal keratinocytes (Figure 4A), as previously demonstrated.¹ These clones seeded onto lethally irradiated feeder layer were characterized by the formation of large and round colonies surrounded by feeder cells, due to the expression of laminin 332 conferring adhesion to extracellular matrix proteins produced by feeder layer.²⁴ In contrast, negative clones maintained the impairment to generate colonies, which is typical of JEB keratinocytes. Restoration of adhesive properties of *LAMB3*-expressing clones was also demonstrated by longer detachment time compared to negative clones. The detachment time for *LAMB3*-expressing clones was comparable to normal keratinocytes (16 min), and negative clones as well as JEB.A649fsX38 cells detached after 5 min of trypsin treatment (Figure 4B). *LAMB3*-expressing clones showed higher proliferation rate compared to negative clones, as assessed *in vitro* by 3-(4,5-dimethylthiazolyl-2)-2,5-diphenyltetrazolium bromide (MTT) assay (Figure 4C). The kinetics of cell growth of laminin-332-expressing clones appeared faster at all time points considered (48, 72, and 96 hr) and was more pronounced at the latest time point.

332 expression *in vivo*, skin equivalent (SE) generated from two clones displaying monoallelic integration of the therapeutic donor (no. 15 and no. 28), one un-corrected clone (no. 71), and untreated JEB.A649fsX38 keratinocytes (data on untreated cells are not shown) were grafted subcutaneously in the back of immunodeficient mice.

Laminin 332 Restoration in a Human-Murine Skin Graft Model

To determine whether *LAMB3*-corrected JEB.A649fsX38 keratinocytes sustain laminin 332 expression *in vivo*, skin equivalent (SE) generated from two clones displaying monoallelic integration of the therapeutic donor (no. 15 and no. 28), one un-corrected clone (no. 71), and untreated JEB.A649fsX38 keratinocytes (data on untreated cells are not shown) were grafted subcutaneously in the back of immunodeficient mice. Four weeks post-transplantation, macroscopic analysis of engrafted areas clearly showed regenerated skin in mice grafted with *LAMB3*-corrected clones (no. 15 and no. 28) and heavily blistered skin in mice grafted with un-corrected clone (no. 71; Figure 5A). Skin biopsies were excised to perform histological, immunofluorescence, and molecular analysis. H&E staining of negative clone-derived SE showed JEB typical blistering phenotype, which was completely reversed in corrected clones. In SE derived from clone no. 71, epidermis (E) completely detached from dermis (D), and SEs derived from clones no. 28 and no. 15 showed continuous dermal-epidermal junction and visible stratification (Figure 5B). Staining using an antibody against human laminin 332 showed strong labeling along the basal membrane in the engrafted SEs derived from genetically corrected clones (no. 15 and no. 28) and undetectable labeling in the SE derived from the negative clone (no. 71; Figure 5C). Genomic DNA from E and D derived from clone no. 71 were included in the molecular assays run on biopsies from *LAMB3*-corrected SE. Targeted integration PCR analyses confirmed the genuine HR of the donor template into Cas9 target site only in biopsies derived from *LAMB3*-corrected SE (Figure 5D). qRT-PCR analysis showed

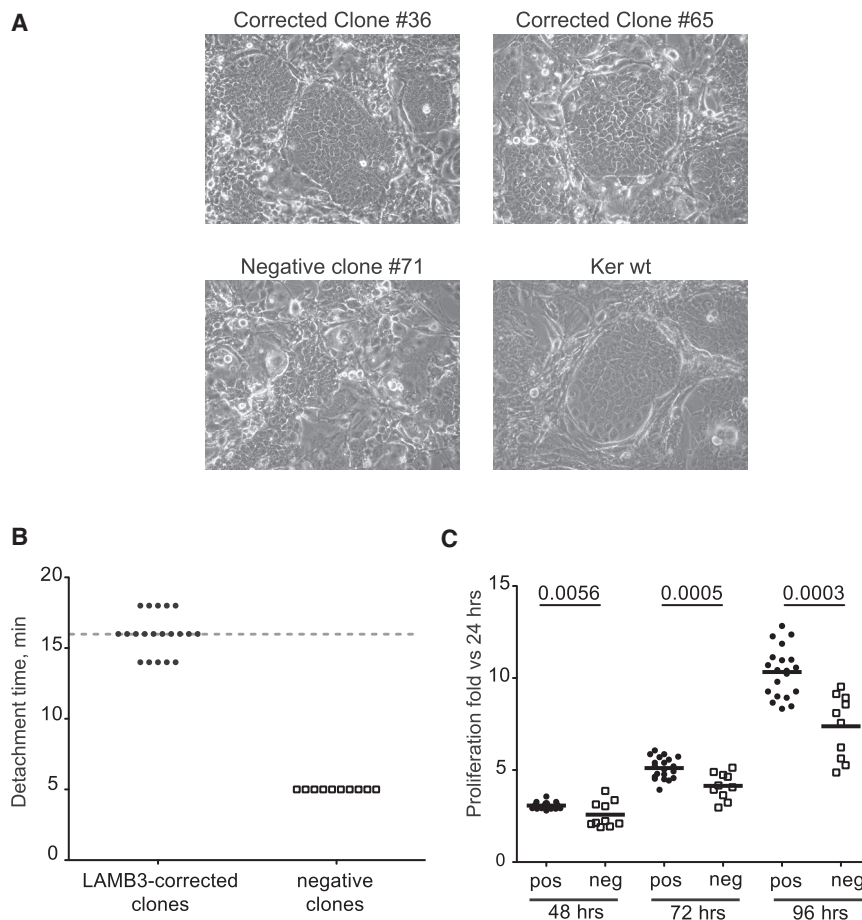


Figure 4. Phenotypic Analysis of LAMB3-Corrected Clones

(A) Microscopic appearance of representative LAMB3-corrected clones (no. 36 and no. 65) grown on feeder layer. As control, negative clone (no. 71) and WT keratinocytes are shown. (B) Detachment time graph indicates the minutes (min) required to 20 LAMB3-corrected clones and 10 negative clones to lose the adhesion properties upon trypsin solution treatment. WT keratinocytes' detachment time is indicated by a dashed line. (C) *In vitro* proliferation of LAMB3-corrected clones (pos) versus negative clones (neg) during time (48, 72, and 96 hr). Results are expressed as fold change from untreated JEB.A649fsX38 keratinocytes at 24 hr, which is set at 1. The bars represent the mean of 3 independent experiments (mean \pm SEM; n = 3). p values are indicated.

more than 20-fold laminin 332 expression in *LAMB3*-corrected SE (Figure 5E) over SE from clone no. 71. These data indicate that monoallelic *in situ* integration of the therapeutic donor sustains sufficient laminin 332 expression to restore the dermal-epidermal junction *in vivo*, in a xenotransplantation model of human SEs on immunodeficient mice.

DISCUSSION

Recently, gene editing approaches were proposed to treat severe diseases, viral infection,²⁵ and monogenic disorders.^{26–35} Any evidence on nuclease-mediated gene editing approach to restore laminin 332 protein, defective in JEB, has not been reported so far. The gene addition strategy based on viral vectors carrying an expression cassette for the correct gene has been the solely available strategy to correct genetic defects for many years. As regards to devastating skin disorders, successful gene therapy protocols based on autologous transplantation of skin keratinocytes transduced with a γ -retroviral vector expressing *LAMB3* or *COL7A1* cDNA were reported to treat JEB^{3,4} and recessive DEB,⁵ respectively. However, gene editing approaches aimed to permanently correct disease-causing mutations or *in situ* integrate therapeutic

cDNA remain an attractive and preferable alternative over the gene addition strategy.

Correction of a single mutation or replacement of quasi-complete gene employs the *in situ* HDR mechanism triggered by nuclease-mediated double-strand breaks (DSBs). A limiting drawback of HDR-mediated correction in clinically relevant cells, e.g., keratinocytes, is represented by the low efficiency of recombination, as reported also by other studies.^{20,34} However, laminin 332 restoration in JEB keratinocytes confers adhesion and proliferation advantages to corrected cells,¹⁶ which could be therefore selected without the need of antibiotic selection

or cell sorting. This observation prompted us to apply CRISPR-mediated HDR to safely *in situ* integrate a therapeutic *LAMB3* cDNA in JEB keratinocytes.

To restore laminin 332 expression in JEB keratinocytes carrying A649fsX38 or R635X mutation in *LAMB3* gene, we employed a first-generation adenoviral vector carrying SpCas9 coupled to a single gRNA tailored to intron 2 of *LAMB3* gene and an IDLV vector bearing the promoterless *LAMB3* cDNA (exon 3–23) flanked by HA. This strategy was designed to potentially address a variety of mutations mapped in *LAMB3* gene,^{36,37} to maintain the transcriptional regulatory elements of the endogenous gene, to optimize laminin 332 expression through a physiological splicing event, and to avoid the activation of neighboring oncogenes if undesired donor integration occurred. The choice of vectors to deliver Cas9 system and the donor template was based on our previous gene editing experience on human keratinocytes.^{20,21} The top-ranked predicted off-targets associated to the designed gRNA did not show presence of indels by Surveyor assay, nevertheless 0.9% indel frequency was detected for OT7 by TIDE, suggesting that genome-wide analysis of off-targets performed by next generation sequencing (NGS)^{38,39} or GUIDE sequencing (GUIDE-seq)⁴⁰ will be necessary before

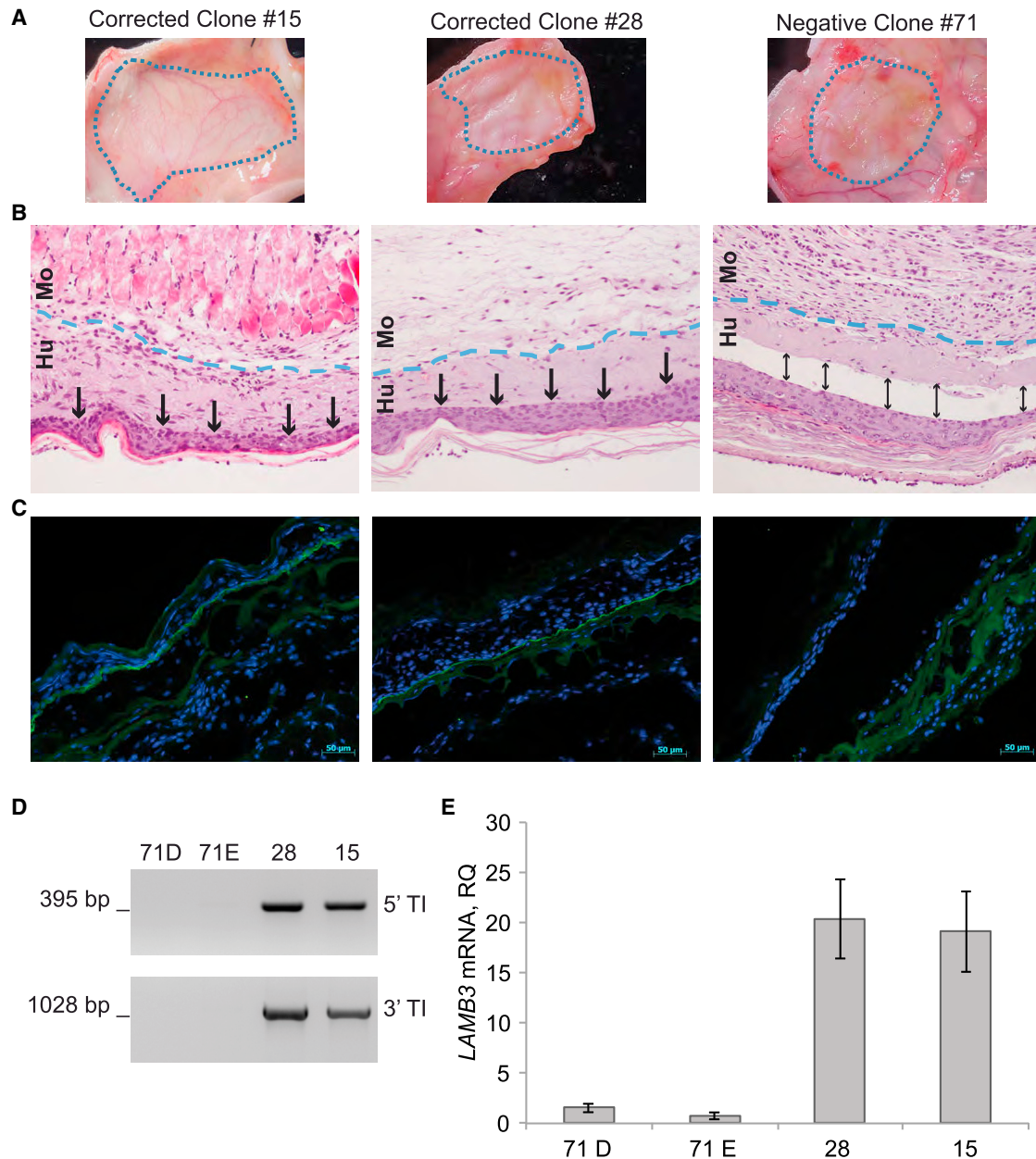


Figure 5. *In Vivo* Skin Regeneration from LAMB3-Corrected Clones

(A) White light epi-illumination of grafts (cyan dotted lines) derived from 2 monoallelic corrected clones (no. 15 and no. 28) and 1 negative clone (no. 71) showed regeneration of human skin transplanted onto immunodeficient mice. (B) H&E staining of skin grafts 4 weeks after transplantation showed normal epidermal differentiation. Cyan dotted lines border the mouse (Mo) and the engrafted human skin (Hu). The arrows indicate BMZ. (C) Immunostaining of laminin 332 confirmed the complete restoration of epidermal-derma junction. The scale bar represents 50 μ m. (D) 5' and 3' TI PCR on genomic DNA extracted from skin biopsies. The size of PCR bands is indicated. (E) *LAMB3* expression in skin biopsies evaluated by qRT-PCR. D, derma; E, epidermis. Data are representative of three independent PCR amplifications (mean \pm SEM; n = 3).

clinical translation. Alternative Cas9 nucleases, e.g., SpCas9-HF1⁴¹ or EvoCas9,⁴² should be considered too.

Genomic analysis of bulk keratinocytes revealed predominant genuine HDR events and rare undesired HDR-independent integration of IDLV-L.B3 donor, as previously observed using similar

IDLV donors^{43,44} (and unpublished data). After 2 rounds of cell detachment, we positively selected cells undergoing a predetermined splicing event, thus expressing the laminin 332 protein that confers adhesive and proliferative advantages. Cells carrying HDR-independent integration of IDLV-L.B3 were negatively selected as demonstrated by genomic analysis of treated JEB bulks and expanded

JEB.A649fsX38 clones. However, these analyses do not rule out the presence of low-frequency (<1%) HDR-independent events in treated JEB keratinocytes, maybe undergoing aberrant splicing affecting *LAMB3* mRNA expression and thereby poor *in vitro* adhesion. 5' TI PCR on single isolated clones showed that 44% (39 out of 88) of expanded clones correctly integrated the donor template by HDR, showing monoallelic correction in the vast majority of them (85%), independently from AdCas9gRNA doses. Based on our previous experience in wild-type keratinocytes²⁰ and on flow cytometric analysis reported in this study, the HDR frequency without selection was <0.5%, thus remarking that selection of gene-corrected cells is needed. Eight clones out of 39 hosted the 5' end of IDLV-L.B3 vector downstream bGH pA signal, further supporting the evidence that HDR-independent integration could be tolerated unless it does not affect *LAMB3* mRNA expression, as demonstrated by RT-PCR analysis. However, the majority of clones screened by 5' TI PCR resulted negative for genuine HDR events, indicating that an optimization of the adhesion-based selection protocol is required. Additional rounds of selection, different detachment time, or lower confluency of the transduced cultures could confer significant improvements to the procedure. Indeed, the experiments performed on JEB.R635X keratinocytes indicated that the adhesion selection protocol set up for JEB.A649fsX38 cells resulted in lower enrichment of *LAMB3*-expressing cells. Analysis at RNA and protein level on isolated JEB.A649fsX38 clones scored laminin 322 expression in all corrected clones. To note, laminin 322 expression level in monoallelic clones, which represent the vast majority of *LAMB3*-corrected clones, was sufficient to recapitulate the adhesive properties and the colony formation typical of normal keratinocytes, as well as their cell growth. Immunodeficient mice were transplanted with SE derived from 2 monoallelic *LAMB3*-corrected JEB.A649fsX38 clones, 1 un-corrected clone, and untreated JEB.A649fsX38 cells. Histological and immunofluorescence analysis showed a completely restored dermal-epidermal junction in biopsies of SE derived from *LAMB3*-corrected clones and a marked and complete detachment of human dermis to epidermis in un-corrected clone and in untreated JEB.A649fsX38 cells. Molecular analyses on DNA and RNA extracted from biopsies confirmed the genuine HR event and the physiological splicing event leading to 20-fold expression of *LAMB3* mRNA over un-corrected or untreated JEB.A649fsX38 cells. Overall, these data provide evidence of the functional correction of laminin 322 defect *in vivo*.

This study represents the first promising application of CRISPR/Cas9-mediated gene editing to restore physiological *LAMB3* expression in keratinocytes derived from JEB patients. The data demonstrated a safe *in situ* integration of the therapeutic *LAMB3* cDNA in JEB keratinocytes, and although the adhesion selection protocol may require further improvement, this novel strategy moves gene editing for JEB closer to the clinic.

MATERIALS AND METHODS

Vectors and Viral Production

For pCCL.LAMB3.HA.SA-cDNALAMB3pA.HA (IDLV-L.B3) construction, *LAMB3* cDNA (3.8 kb in length), spanning from exon 3

to 23 and including the splice acceptor of exon 3, was amplified from MFG-LAMB3.³ Promoterless *LAMB3* cDNA was cloned upstream of bGH pA signal, between 5' and 3' HA (90 and 413 bp in length, respectively), both from intron 2 of *LAMB3* gene. The donor cassette was finally cloned in reverse orientation into pCCL.LV.poly-linker vector,⁴⁵ generating the transfer plasmid to be packaged into integration defective lentiviral vector (IDLV-L.B3).

For AdCas9gRNA construction, pShuttle.CRISPR.gRNA1LAMB3 was generated by cloning fragment U6.gRNA1LAMB3.CBH.hSpCas9.bGH.pA from plasmid pX330.gRNA1.LAMB3 in pShuttle from AdEasy Human Adenoviral Vector System (Stratagene). Briefly, pX330.gRNA1.LAMB3 was obtained by cloning gRNA1LAMB3 in pX330-U6-Chimeric_BB-CBH-hSpCas9 plasmid (Addgene no. 42230) by oligo annealing into *BbsI* sites (<http://www.addgene.org>). Then, pX330.gRNA1.LAMB3 was digested with *PciI* and *NotI*, and fragment U6.gRNA1Lamβ3.CBH.hSpCas9.bGH.pA was cloned in pShuttle from AdEasy (AdEasy Adenoviral Vector System; Stratagene), previously digested with *NotI* and filled in. pShuttle.CRISPR.gRNA1LAMB3 was linearized with *PmeI*, de-phosphorylated, and co-transformed with pAdEasy-1 from AdEasy system in BJ5183 bacteria to produce Ad5 genomic plasmid Ad.CRISPR.LAMB3gRNA1.

IDLV viral stocks pseudotyped with vesicular stomatitis virus G protein were prepared by transient co-transfection of HEK293T cells with transfer vector, pMD.Lg/pRRE.D64VInt packaging plasmid, pMD2.VSV-G envelope-encoding plasmid, and pRSV-Rev.²⁰ Vector particles were concentrated by ultracentrifugation and titrated by Taqman RT-PCR with PCR primers and 6-carboxyfluorescein (FAM) probe for WPRE (Applied Biosystems). IDLV-L.B3 titer was 10⁹ vg/mL by qPCR.

AdCas9gRNA vector was produced and titrated as previously described.⁴⁶ Adenoviral titer was 1.9 × 10¹¹ vg/mL by qPCR. The MOI reported for this virus indicates the amount of vg per cell.

Cell Culture

Primary keratinocytes from JEB patient carrying the homozygous *LAMB3* mutation in exon 14 (c.1945dupG)¹⁷ were immortalized to facilitate clonal analysis. Early-passage primary JEB keratinocytes were infected with supernatants of pLXSN-based retroviral vector containing cDNA sequence of E6 and E7 proteins from papilloma virus type 16⁴⁷ and referred to as JEB.A649fsX38 keratinocytes. The karyotype of JEB.A649fsX38 keratinocytes was analyzed (four metaphases) as described in Chamorro et al.⁴⁷ and showed aneuploidy with redundant trisomy for chromosomes 9 and 20. Chromosome 1, which hosts *LAMB3* gene, was diploid in 50% of analyzed metaphases, although in other metaphases, we detected an extra chromosome 1 (Figure S11). The second immortalized JEB keratinocytes line (JEB.R635X) used in this study was established as described in Hammersen et al.¹⁸ Feeder layer was required for JEB keratinocyte cell growth; thus, they were plated onto lethally irradiated 3T3-J2 cells. JEB.A649fsX38 keratinocytes were cultured in keratinocyte growth medium (cFAD medium and epidermal growth factor; KC medium),

diluted 1:1 with Cnt-57 (progenitor cell-targeted [PCT] epidermal keratinocyte medium low bovine pituitary extract [BPE]; CellnTEC Advanced Cell Systems AG, Bern, Switzerland). JEB.R635X keratinocytes were cultured in keratinocyte growth medium (cFAD medium and epidermal growth factor; KC medium), diluted 1:1 with KGM-Gold (Lonza; Euroclone, Milan, Italy). Mouse 3T3-J2 cells⁴⁸ were grown in DMEM supplemented with 10% donor bovine serum (Gibco; Thermo Fisher Scientific, Monza, Italy), 50 IU/mL penicillin-streptomycin, and 4 mmol/L glutamine.

Transduction of JEB Keratinocytes, Adhesion Selection, and Isolation of Single-Cell Clones

For editing analysis, JEB cells were plated 24 hr before transduction (without feeder layer) and then were transduced with AdCas9gRNA in serum-free medium containing viral particles at MOIs 10^4 , 2×10^4 , 5×10^4 , or 10^5 vg/cell. After 1 hr at 37°C, the viral mixture was replaced with Cnt-57 + KC medium. For HR-mediated correction, JEB keratinocytes (2.5×10^5) were first infected with IDLV-LB3 (4.32×10^2 vg/cell) in suspension, in the presence of polybrene (8 µg/mL), and seeded in 6-well plates onto lethally irradiated 3T3-J2 cells. After 6 hr, Cnt-57 medium was added to cells. After 24 hr, AdCas9gRNA transduction was performed in serum-free medium containing viral particles at MOIs 10^4 , 2×10^4 , 5×10^4 , or 10^5 vg/cell. After 1 hr at 37°C, the infection mixture was replaced with Cnt-57 + KC medium.

To enrich *LAMB3*-corrected cells, 4 days post-transduction JEB keratinocytes transduced with IDLV-LB3 (4.32×10^2 vg/cell) and AdCas9gRNA (MOIs 10^4 and 2×10^4 vg/cell) were washed with PBS and incubated with trypsin solution at 37°C. Cells that detached after 4 min of incubation were removed and referred as un-corrected (negative) JEB cells. Adherent cells were further incubated with trypsin solution at 37°C for additional 4 min and named *LAMB3*-corrected JEB keratinocytes. After trypsin treatment, cells were plated onto lethally irradiated 3T3-J2 cells in KC medium diluted 1:1 with Cnt-57. To isolate single-cell clones, JEB.A649fsX38 cells transduced with IDLV-LB3 (4.32×10^2 vg/cell) and AdCas9gRNA (MOIs 10^4 and 2×10^4 vg/cell) were limiting diluted, plated onto lethally irradiated 3T3-J2 cells, and cultured in KC medium diluted 1:1 with Cnt-57.

Editing Efficiency Analysis and DNA Sequence Analysis

To analyze editing efficiency and specificity, genomic DNA was extracted using DNeasy MiniKit (QIAGEN, Milano, Italy). To evaluate on-target editing, gRNA target region from JEB cells transduced at MOI 10^4 , 2×10^4 , 5×10^4 , and 10^5 vg/cell or un-transduced (mock transduced) was amplified by PCR (Platinum Taq High Fidelity; Thermo Fisher Scientific, Monza, Italy), using primers Cell1Lamb3for and Cell1Lamb3rev (Table S1), and analyzed with TIDE software to evaluate indels frequency.

To amplify the off-targets predicted by two public webservers (<http://crispr.mit.edu> and CRISPR direct [<https://crispr.dbcls.jp>]), we designed several primers listed in Table S1. The amplification products were analyzed by Surveyor assay, according to SURVEYOR Mutation

Detection Kit (Transgenomic, Omaha, NE) manufacturer's protocol. Briefly, 72 hr after transduction, PCR amplifications were performed on genomic DNA and then PCR products were denatured, digested by Cel-I nuclease, and finally subjected to 2% agarose gel electrophoresis. 100 bp DNA ladder from New England Biolabs (Massachusetts, USA) was used as molecular weight marker. For DNA sequence analysis of on-target indels, PCR products were subcloned into a PCR2.1 TOPO vector (Thermo Fisher Scientific, Monza, Italy) and Sanger sequenced (Eurofin s.r.l Vimodrone, Italy). Sanger sequences were analyzed accordingly to a bioinformatics pipeline (D.B., V. Marigo, and A.R., unpublished data) to accurately evaluate the percentage and type of indels upon AdCas9gRNA treatment.

PCR and Semiquantitative and qRT-PCR Analyses

Genomic DNA was isolated from JEB cells or skin biopsies with a QIAmp DNA Mini kit (QIAGEN, S.p.A., Milan, Italy), according to manufacturer's protocols. To analyze correct integration of the donor in *LAMB3* locus, we performed target integration PCRs (PCR GoTaq, Promega, Milan Italy) on genomic DNA using primers TI5'Lamb3for and TI5'Lamb3rev to analyze the correct integration at 5' region and primers TI3'Lamb3for and TI3'Lamb3rev to analyze the correct integration at 3' region (see Table S2 for primers list). PCR cycles for TI5' are 95°C 30 s, 57°C 30 s, and 72°C 30 s; PCR cycles for TI3' are 95°C 30 s, 55°C 30 s, and 72°C 1 min. PCR products were loaded on 1% agarose gel, and 1 kb or 100 bp DNA ladders from New England Biolabs (Massachusetts, USA) were used as molecular weight markers.

To investigate the integration of the 5' end of IDLV vector, we performed modified target integration 3' PCR (TaKaRa LA Taq, USA) on genomic DNA using primers TI3'Lamb3for and Lamb3-Ex3-RC. PCR cycles are as follows: 98°C 30 s; 60°C 30 s; and 68°C 2 min. For allelic discrimination, a three-primer-based PCR (TaKaRa LA Taq, USA) was performed to discriminate between parental allele (TI5'Lamb3for and TI3'Lamb3rev) and donor-integrated allele (TI5'Lamb3for and TI5'Lamb3rev). PCR cycles are as follows: 98°C 20 s; 55°C 30 s; and 68°C 5 min.

Total RNA from JEB cells or skin biopsies was isolated with RNeasy Mini kit plus (QIAGEN, Milan, Italy), according to manufacturer's protocols. cDNA from JEB cells was synthesized in a 20 µL reaction using 200 ng total RNA and ImProm-II Reverse Transcription System (Promega, Milan, Italy). cDNA from skin biopsies was synthesized in a 20-µL reaction using 400 ng and SuperScript III (Life Technologies, Monza, Italy).

To evaluate *LAMB3* mRNA expression, semiquantitative RT-PCR was performed with primers Lamb3Exon13_F and Lamb3Exon15_R. PCR cycles are as follows: 95°C 30 s; 60°C 30 s; and 72°C 45 s. As housekeeping control, GAPDH mRNA was amplified with primers GAPDH-F and GAPDH-R. PCR cycles are as follows: 95°C 30 s; 60°C 30 s; and 72°C 45 s.

qRT-PCR was performed in 96-well plates on ABI Prism 7900 Sequence Detection System (Applied Biosystems, Monza, Italy)

with TaqMan Universal PCR Master Mix and Taqman Gene Expression assay (Applied Biosystems), with a final reaction of 25 μ L. PCR primers and 6-carboxyfluorescein (FAM) probes for *GAPDH* (Hs99999905_m1) or *LAMB3* (Hs00989722_m1) were purchased from Applied Biosystems. Reactions were performed according to manufacturer's instructions. The relative expression was normalized to the level of *GAPDH* in the same cDNA using $2^{-\Delta\Delta C_t}$ quantification. The mean of replicated relative quantity (RQ) values for each sample was calculated.

Flow Cytometry

For flow cytometric analysis of LAMB3 expression, after washing with PBS + 5% fetal bovine serum (FBS) + 2.5 mM EDTA, JEB cells were stained with antigen-presenting cell (APC)-conjugated anti-feeder antibody (Miltenyi Biotec, Bologna, Italy) accordingly to manufacturer's protocol. Cells were fixed and permeabilized using the Fix and Perm cell permeabilization reagents kit (Invitrogen, Monza, Italy), according to manufacturer's protocol, before intracellular staining of LAMB3 with anti- β 3 chain monoclonal antibody K140 (diluted 1:30; a kind gift from Prof. Castiglia, Istituto Dermopatico dell'Immacolata, IRCCS, Rome, Italy). As secondary antibody, Alexa 488 donkey anti-mouse antibody (Life Technologies, Monza, Italy; 1:1,000) was used. Fluorescence was acquired on a FACS CANTO cytometer (BD Biosciences, Milan, Italy). Intracellular LAMB3 expression was evaluated after feeder exclusion (APC- cells). The percentage of LAMB3-positive cells was determined after subtracting the percentage of LAMB3-positive events in untreated cells. All flow cytometry experiments were analyzed with the FlowJo software (Tree Star).

Western Blotting

Cell lysates were extracted with radioimmunoprecipitation assay (RIPA) buffer containing proteinase inhibitor cocktail (Complete Mini, EDTA-free; Roche Diagnostics, Monza, Italy), incubated for 30 min on ice and centrifuged at 14,000 rpm for 30 min. The amount of proteins was determined using Bradford assay (Bio-Rad, Milan, Italy). Sixty micrograms of JEB keratinocytes extracts and 10 μ g of primary keratinocyte extracts were separated on 4%–12% gradient NuPAGE Bis-Tris Gel (Life Technologies, Monza, Italy) under reducing condition. Western blots were performed with a mouse monoclonal antibody against human Kalinin B1 (Kalinin B; 1:1,000; BD Transduction Laboratories, Milan, Italy). Kalinin B1 is an alternative name of laminin subunit beta-3 protein coded by *LAMB3* gene according to Uniprot (<https://www.uniprot.org/uniprot/Q13751>). For protein loading normalization, a mouse monoclonal antibody anti- β -actin (sc-81178; 1:1,000; Santa Cruz Biotechnology, Heidelberg, Germany) was used. Horseradish-peroxidase-conjugated anti-mouse antibody (diluted 1:5,000) was used for chemiluminescent detection (Amersham; GE Healthcare Europe, Milan, Italy).

Keratinocyte Growth and Detachment Analysis

JEB clone proliferation was measured by MTT reduction colorimetric assay (Sigma-Aldrich, Milan, Italy). For the time course analysis, 4×10^4 cells/well were plated without feeder layer in KC medium

diluted 1:1 with CnT-57, a well of a 24-well plate per time point, and incubated at 37°C for 24, 48, 72, or 96 hr. At each time point, cells were incubated with 500 ng/mL MTT solution at 37°C for 90 min, and then MTT solution was gently removed. The incorporated dye was solubilized in acidified isopropanol and determined by absorbance measurement (A_{570nm}). Results are expressed as fold change from untreated control cells at 24 hr, which is set at 1 and as mean of 3 independent experiments.

Detachment time analysis was performed by measuring the time of incubation with trypsin solution necessary to all cells of a sub-confluent culture to lose adhesive properties.

Bioengineered Skin Preparation and Grafting on Immunodeficient Mice

The SE or bioengineered skin corresponds to a 3D culture resembling the basic architecture of skin. It comprises a dermal hydrogel matrix (fibrin) containing embedded live human fibroblasts and an epidermal component formed by human keratinocytes (genetically corrected or non-corrected) added to the dermal matrix and grown until confluence (keratinocyte layer). Bioengineered skin grafts were prepared from 4 samples: 2 *LAMB3*-corrected clones; 1 negative clone; and untreated JEB.A649fsX38 cells as previously described.⁴⁹ Bioengineered SEs were grafted subcutaneously in the back of immunodeficient nu/nu mice ($n = 3$ /sample) according to Barrandon et al.⁴⁹ Four weeks after grafting, mice were sacrificed and grafts harvested for skin immunohistochemistry and molecular analysis. Animal studies were approved by our institutional animal care and use committee according to all legal regulations.

Immunofluorescence and Immunohistochemistry

Immunofluorescence analysis for laminin 332 expression on skin biopsies from mice was performed on 7- μ m sections. After blocking with 10% fetal bovine serum + 3% BSA in PBS, sections were incubated with 50 μ g/mL unconjugated Fab anti-immunoglobulin G (IgG) antibody. After wash, sections were incubated with mouse monoclonal GB3 antibody (diluted 1:5; a kind gift from Prof. Castiglia, Istituto Dermopatico dell'Immacolata, IRCCS, Rome, Italy). As secondary antibody, Alexa 488 donkey anti-mouse antibody (Life Technologies; 1:1,000) was used. Staining with 0.1 μ g/mL DAPI was performed for nuclear staining. Slides were mounted with Dako Fluorescent Mounting Medium (Dako Italia SRL, Milan, Italy) and analyzed with a Zeiss Axioskop 40 FL fluorescence microscope equipped with a digital camera AxioCam and AxioVisionRel version 4.8 software for image processing (Carl Zeiss, Milan, Italy).

Histological samples of the whole mouse-human skin "sandwich" were formalin fixed and H&E stained as described in Chamorro et al.⁴⁷

Statistical Analysis

Two-way ANOVA or Student's t test was performed for statistical significance. All values in each group were expressed as mean \pm SEM.

SUPPLEMENTAL INFORMATION

Supplemental Information includes eleven figures and two tables and can be found with this article online at <https://doi.org/10.1016/j.ymthe.2018.07.024>.

AUTHOR CONTRIBUTIONS

D.B., F.M., and F.C. designed experiments; D.B. performed experiments *in vitro* and *ex vivo* on skin biopsies and analyzed data; F.M., F.C., C.P., and S.B. performed experiments *in vitro*; C.H. provided immortalized JEB carrying the R635X mutation; M.C., M.D.R., and F.L. were involved in SE production and grafting experiments; and S.C. and V.A. packaged the first generation AdCas9gRNA vector. D.B. and A.R. wrote the manuscript.

CONFLICTS OF INTEREST

The authors have no conflicts of interest to disclose.

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