Generation of High-Titer Self-Inactivated γ-Retroviral Vector Producer Cells

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INTRODUCTION

The γ-retroviral vector is a gene delivery vehicle that is commonly used in gene therapy. Despite its efficacy, its strong enhancers contributed to malignant transformations in some hematopoietic stem cell (HSC) gene therapy trials. A safer version without viral enhancers (SIN) is available, but its production is cumbersome, as high titers can only be obtained in transient transfection. Our aim was to develop a system that could easily generate high-titer SIN vectors from stable producer cells. The use of the cytomegalovirus enhancer-promoter sequence to generate the full-length genomic RNA combined to producer cells. The use of the cytomegalovirus enhancer-promoter sequence to generate the full-length genomic RNA combined to sequences that decrease transcriptional readthrough (WPRE and strong polyadenylation sequences) led to 6 × 10^6 infectious units (IU)/mL of a SIN GFP vector in transient transfection. The incorporation of a blasticidin selection cassette to the retroviral plasmid allowed the generation of stable clones in the 293Vec packaging cells that release 2 × 10^7 IU/mL and 1.4 × 10^7 IU/mL of a SIN GFP and a SIN PIGA vector, respectively. A titer of 1.8 × 10^6 IU/mL was obtained with a SIN vector containing the long 8.9-kb COL7A1 cDNA. Thus, an efficient process was established for the generation of stable 293Vec-derived retrovirus producer cells that release high-titer SIN vectors.

The γ-retroviral vector can be produced by transient transfections or with stable packaging cell lines. The latest option is preferable, as the manufacturing is easier, more reproducible, less expensive, and more suitable for large-scale productions needed for late-phase trials and commercialization.

The γ-retroviral vector is widely used in ex vivo gene therapy applications for the treatment of genetic diseases and cancer. Successful examples of therapies using this vector are Strimvelis, Yescarta, and Zalmoxis, which have been recently approved for commercialization.3–5 Strimvelis is a HSC gene therapy treatment for adenosine deaminase (ADA) deficiency,6 and Yescarta is a treatment for non-Hodgkin lymphoma in which the patient’s own T cells are genetically modified to express a chimeric antigen receptor directed toward CD19.7 Zalmoxis is a gene therapy treatment that controls graft versus host disease (GVHD) in patients undergoing haploidentical bone marrow transplantation. In this approach, autologous T cells are engineered in vitro to express the herpes simplex virus thymidine kinase (HSV-TK) gene. After grafting, these cells would be eliminated in the presence of the nucleoside analog ganciclovir if the patient develops GVHD.7

In 2000, Cavazzano-Calvo and colleagues used a γ-retroviral vector to successfully treat X-linked severe combined immunodeficiency (SCID-X1) patients.8 Mutations in the interleukin-2 receptor γC (IL2RG) gene in SCID-X1 lead to an absence of T cell and natural killer cell development and B cell dysfunction.9 In the gene therapy trial, harvested HSCs were transduced in vitro with a γ-retroviral vector containing the IL2RG cDNA. Cells were subsequently reinfused in patients, and a normal hematopoiesis with a full immune reconstitution was observed shortly after in the majority of patients.10 Unfortunately, 4 patients in the original French trial, as well as another one in a similar trial in London, developed T cell acute lymphoblastic leukemia between 2 and 6 years after gene therapy.11–13 Other cases of leukemias and myelodysplastic syndromes were reported in HSC γ-retroviral gene therapy for Wiskott-Aldrich syndrome and chronic granulomatous disease.14–17 In all these patients, the integration site of the vector was near proto-oncogene sequences like PRDM16, SETBP1, and MDS1-EVI1 that became transcriptionally activated, owing to the strong viral enhancers present in the vector. In addition, several chromosomal alterations were observed in the

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leukemic cells of these patients. Surprisingly, no leukemia cases have been reported in HSC gene therapy for ADA, suggesting that other factors, like the nature of the transgene or of the disease treated could have also contributed to the malignant transformation mediated by the γ-retroviral vector. Therefore, several groups have clearly shown in vitro and in animal models that a self-inactivated (SIN) γ-retroviral or lentiviral vector that has its enhancers deleted was less oncogenic than its wild-type counterpart, as long as it contains a relatively weak internal promoter. Based on these results, a new SCID-X1 gene therapy clinical trial was launched with a SIN γ-retroviral vector. More than 6 years after treatment, there were no leukemia cases reported in the 7 patients who have been followed, and the efficacy was comparable to that of previous trials in which a regular γ-retroviral vector was used. 

Like the γ-retroviral vector, the lentiviral vector is integrative, and it is used in HSC gene therapy. The SIN lentiviral vector has a good safety profile with, so far, no oncogenic events reported in HSC clinical trials. The toxicity of some lentiviral proteins and the vesicular stomatitis virus-G (VSV-G) envelope that is commonly used to pseudotype the vector limits the manufacturing to transient systems. The SIN γ-retroviral vector manufactured for the SCID-X1 trial was pseudotyped with the non-toxic envelope from the gibbon ape leukemia virus (Galv); but it was still produced by transient transfections in order to get relatively good titers, around 10^6 infectious units per milliliter (IU/mL).

A few years ago, we constructed three retrovirus packaging cell lines that release high-titer viral particles pseudotyped with the amphotropic (293Vec-Ampho), the RD114 (293Vec-RD114), and the Galv (293Vec-Galv) envelopes. These cell lines can produce regular γ-retroviral vectors with titers in the range of 10^7 IU/mL, and they have the properties to grow in suspension and serum-free media. In this study, our goal was to design an optimized SIN vector that could lead to high-titer vectors with the 293Vec packaging cell lines. The SIN γ-retroviral vector that was constructed led to high-titer vector production in transient transfection. High-titer retrovirus producer clones were also easily established from the 293Vec-RD114 and 293Vec-Ampho cells with three different SIN retroviral vectors containing the GFP gene and two therapeutic transgenes.

RESULTS
Optimization of a SIN γ-Retroviral Vector
The introduction of strong enhancer-promoter sequences in place of U3 in the 5' long terminal repeat (5'LTR) sequence has been proposed to increase titers of SIN γ-retroviral and lentiviral vectors. In this study, the enhancer-promoter sequences of the cytomegalovirus (CMV) and of the Rous sarcoma virus (RSV) were compared in the context of a SIN MFG-derived γ-retroviral vector. The synthetic sequence composed of the CMV enhancer sequence fused to the chicken β-actin promoter (CAG) was also tested, as its strength is superior to that of the CMV and the RSV enhancer-promoter sequences in several eukaryotic cell lines. Vector optimization was conducted with the G418 gene under the control of the human elongation factor-1 α promoter (EF-1α), and their designs are displayed in Figure 1.

Transient productions of SIN.RSV generated a titer of 3 × 10^5 IU/mL, and a 5.6-fold higher titer was obtained with the SIN.CMV vector. The titer of the SIN.CAG vector that was also tested was intermediary at 9.6 × 10^5 IU/mL. The woodchuck hepatitis posttranscriptional regulatory element (WPRE) sequence and strong polyadenylation (poly(A)) sequences were next incorporated to the SIN.CMV vector to counteract the readthrough of the MLV poly(A). WPRE increased the titer from 1.7 × 10^6 to 4.2 × 10^6 IU/mL, and a titer of 6 × 10^6 IU/mL was obtained with the incorporation of the simian virus 40 (SV40) poly(A) in the R region. The addition of the bovine growth hormone (BGH) poly(A) downstream to the 3'LTR of the SIN.CMV.W.SV gave a titer of 6.2 × 10^6 IU/mL (Figure 2). These results showed that the CMV sequence is superior to the RSV and CAG sequences for the production of the MFG-derived SIN γ-retroviral vector, and that the subsequent additions of WPRE and poly(A) sequences have a positive effect on viral production.

Production of SIN GFP γ-Retroviral Vectors in Stable Packaging Cell Lines
Vectors containing the CMV enhancer-promoter sequences were next tested during stable viral productions. The 293Vec-RD114 packaging cells were co-transfected with each construct and a hygromycin resistance (Hyg') plasmid, and viral titers from supernatants of sorted GFP-positive cell populations were measured. Results were
SIN γ-Retroviral Vectors Containing Therapeutic Transgenes

The performance of the SIN γ-retroviral vector was then assessed with 2 different therapeutic transgenes. The first one is the 8.9-kb COL7A1 cDNA that could be used to correct keratinocytes and fibroblasts of recessive dystrophic epidermolysis bullosa (RDEB) patients. A vector with such a long transgene would be less efficiently packaged and would lead to low titers. We hypothesized that the benefit of WPRE could be cancelled by its size (700 bp), which would further decrease the packaging ability of a COL7A1 vector. Two SIN vectors with or without WPRE were then constructed, and their titers were compared in transient productions. Similar vectors were also constructed with a smaller therapeutic transgene: the COL7A1 vector. For simplicity, SIN vector final versions used in the study will be referred to hereinafter as SINVec.GFP, SINVec.PIGA, and SINVec.COL7A1 (-W).

An Efficient Selection Method for the Stable Production of SIN Vectors

A blasticidin resistance (BSD') cassette was introduced in the SIN GFP retroviral plasmid to facilitate the generation of stable retrovirus producer cells (Figure S2). The BSD' gene was placed under the control of a minimal promoter fragment from the HSV-TK gene, and this cassette was ligated to the retroviral plasmid. Our hypothesis was that several copies of vectors would be needed to confer blasticidin resistance; this would lead to more integrated plasmid copies in positive cells as well as cells that release vector at higher titers. To validate our hypothesis, packaging cells were transected with the GFP SIN plasmid containing the blasticidin cassette or co-transfected with the GFP SIN plasmid and a Hygro' plasmid for comparison. Cells selected with blasticidin were 91% GFP positive, while only 48% of the cells were fluorescent after hygromycin selection (Figure S3). Furthermore, sorted GFP-positive cells were 2.1-fold more fluorescent if cells had been selected in blasticidin versus hygromycin. There were also 9.9 plasmid copies integrated per cell in the BSD' cell population, while only 2.9 copies were present in the Hygro' cells. These differences between the two cell populations led to the production of vectors at 7 × 10^6 IU/mL from the Hygro' cells and 1.7 × 10^6 IU/mL from the BSD' cells (Table 1). Thus, the addition of a stringent blasticidin selection cassette to our SIN retroviral plasmid was an efficient strategy to select high-titer retrovirus producer cells.

Generation of Stable Retrovirus Producer Clones with SIN Vectors

SIN vectors with GFP, COL7A1, and PIGA were next tested for stable vector productions in packaging cell clones. WPRE was kept for the SIN GFP and PIGA vectors but not for the COL7A1 vector, as this sequence had no positive effect on titers (Figure 4). The blasticidin cassette was introduced in each of these retroviral plasmids that were transfected in the 293Vec-RD114 packaging cells for the GFP and the PIGA vectors and in the 293Vec-Ampho packaging cells for the COL7A1 vector, as amphotropic pseudotyped vectors are more efficient for transducing human fibroblasts. All the 43 clones screened with the GFP vector had titers above 2 × 10^6 IU/mL, and for the PIGA vector, 90% had titers above 6 × 10^5 IU/mL. For the COL7A1 vector, 80% of the clones were above 2 × 10^5 IU/mL, with an average titer of 6 × 10^5 IU/mL (Figure 5A). Vectors produced from the best clones generated titers of 2.6 × 10^7 IU/mL for the GFP vector, 1.4 × 10^7 IU/mL for the PIGA vector, and 1.8 × 10^6 IU/mL for the COL7A1 vector (Figure 5B). These packaging cell clones had an average of 6.5 plasmid copies per cell and demonstrated a stable viral vector production for a 3-month period.
Transduction of Primary Cells with SIN Vectors

We next tested the transduction efficacy of the SIN GFP and COL7A1 vectors on primary cells. Human primary T lymphocytes were activated and transduced with the SIN GFP vector produced from the best 293Vec-RD114 clone in the presence of retronectin. A high transduction efficiency of 70% was achieved at a MOI of 5 (Figure 6A). Next, fibroblasts and keratinocytes from one RDEB patient negative for collagen VII were transduced with virus produced from our best 293Vec-Ampho clone. A strong transduction was achieved, as keratinocytes and fibroblasts were 74% and 77% positive for collagen VII expression, respectively (Figure 6B). We could conclude that SIN vectors produced from stable 293Vec clones have the potential to be used in clinical gene therapy trials.

DISCUSSION

Retroviral and lentiviral vectors are commonly used in ex vivo approaches that involve the genetic modification of HSCs and T lymphocytes. Although these vectors have been extremely safe when used with T cells, leukemias and myelodysplastic syndroms have been reported in some HSC gene therapy clinical trials with the γ-retroviral vector.\(^\text{11,15,17,18}\) The absence of genotoxicity with the lentiviral vector is mainly due to its SIN design that also leads to a good safety profile when it is applied to the γ-retroviral vector.\(^\text{20,22}\) The main objective of this study was to design a SIN γ-retroviral plasmid that could easily generate high-titer stable retrovirus producer cells, a feature that is currently not achievable with the lentiviral vector.

In this study, several modifications introduced in the MFG-derived γ-retroviral plasmid to increase viral titers were tested in transient transfections. The CMV enhancer-promoter sequence in place of U3 in the 5′ LTR and the addition of the WPRE sequence and two strong poly(A) sequences increased the titer of a SIN GFP vector up to \(6 \times 10^6\) IU/mL. The addition of a BSD\(^\circ\) cassette in the SIN plasmid allowed the easy selection of high-titer stable virus producer cells with the 293Vec-RD114 and 293Vec-Ampho packaging cells. Titers of \(2.6 \times 10^7\) and \(1.4 \times 10^7\) IU/mL were obtained with retrovirus producer cell clones with the SINVec.GFP and the SINVec.PIG-A vectors, respectively. SINVec.COL7A1 (-W) was produced at a titer of \(1.8 \times 10^6\) IU/mL.

It has been well established that internal promoters in SIN γ-retroviral vectors interfere with the activity of the 5′ LTR. This promoter competition limits the production of the genomic RNA that leads to low vector titers.\(^\text{35}\) In addition, the deletion of U3 in the 3′ LTR of the SIN γ-retroviral vector leads to a defect in genomic RNA exportation.\(^\text{37,48}\) The use of strong enhancer-promoter sequences in place of the U3 region in the 5′ LTR is a strategy that has been proposed to overcome this limitation with γ-retroviral as well as with lentiviral vectors.\(^\text{35–37}\) It has been previously shown with both vectors that the RSV LTR sequence led to higher titers than the CMV enhancer-promoter sequence.\(^\text{35,36}\) In this study, the results were opposite as a 6-fold higher titer was found with the SIN.CMV vector versus the SIN.RSV vector in transient transfections (Figure 2). This difference could be explained by different vector backbones used among the studies: lentiviral from HIV type 1,\(^\text{36}\) γ-retroviral derived from the friend spleen focus-forming virus,\(^\text{35}\) and γ-retroviral derived from MLV (our study). Small nucleotide differences in the enhancer-promoter sequences used among the studies could also explain this discrepancy.

The WPRE sequence is often incorporated in lentiviral and retroviral vectors to increase viral titers as well as transgene expression in transduced cells.\(^\text{49–51}\) Its effect has been mainly attributed to a decrease from transcriptional readthrough that occurs with the relatively weak retrovirus poly(A) sites.\(^\text{10,52}\) In our study, WPRE increased by 3-fold the titer of a SIN GFP vector produced by transient transfections, and this effect was 10-fold more pronounced when vectors were produced from stable packaging cell lines (Figures 2 and 3). This difference could be due to the limited amount of genomic RNAs available in the stable packaging cell lines that could be strongly increased by the addition of WPRE. On the contrary, the amount of genomic RNAs should not be that limiting in transient viral productions, as large quantities of plasmids would be present in the transected cells. A similar trend has been observed with the incorporations of the two poly(A) signals: significant increases in titers were only obtained with stable retrovirus producer cells (Figure 3).
It has been shown that retroviral vectors that exceed the wild-type proviral RNA in size are produced at lower titers. The 10-fold lower titers obtained in transient transfections with the SIN Vec.COL7A1 or the SIN Vec.COL7A1 (-W) vector versus the SIN Vec.GFP or the SIN Vec.PIGA vector were then expected (Figures 2 and 4). Indeed, the size of the SIN Vec.COL7A1 (-W) vector leads to a genomic RNA of 10.2 kb that exceeds by 1.9 kb the size of the wild-type genomic RNA. The addition of WPRE did not increase the titer of the SIN Vec.COL7A1 vector, but its effect was observed with the GFP and the PIGA vectors with a 2.5-fold and a 2.9-fold titer increase, respectively (Figures 2 and 4). One logical explanation for this result could be that the positive effect of WPRE is cancelled by its size, which would further decrease the packaging ability of the SIN COL7A1 vector.

Sorted GFP packaging cells co-transfected with the SIN.CMV.W. SV-BGH plasmid and a Hygro plasmid produced vectors with titers of \(7 \times 10^6\) IU/mL (Figure 3). We hypothesized that the genomic RNAs were still limiting, because the 293Vec packaging cell line produces less virus than PG13 cells.33 In the second study, the SIN vector was introduced in the packaging cells by recombinase-mediated cassette exchange in a predefined locus of elevated transcription activity. With this strategy, SIN vector titers of \(2 \times 10^7\) IU/mL and \(4 \times 10^6\) IU/mL were obtained with a T cell receptor and a COL7A1 cDNA transgene, respectively. These titers are very similar to those found in our study. The advantage of this strategy is that only one copy of the vector is present in the packaging cell, which would facilitate its characterization. However, this strategy is cumbersome, as two cloning steps were necessary to obtain the final vector producer cells. The first cloning was for the exchange of the vector, and the second one was to remove the residual sequences that were required for targeting and selection.57 The 293Vec cells were also used in this study, suggesting that the potency of this packaging cell line to release high-titer vectors is key for the production of SIN vectors.

In conclusion, this work showed that high-titer stable producer cells could be easily established with an optimized SIN \(\gamma\)-retroviral vector in 293Vec cells. We conclude that the stable production of SIN \(\gamma\)-retroviral vectors with the 293Vec platform will be a more efficient, a more reliable, and a cheaper option for late-stage trials and commercialization than the production of SIN \(\gamma\)-retroviral or lentiviral vectors in transient transfection.

MATERIALS AND METHODS

Ethics

The study was approved by the ethics committee of the CHU de Québec - Université Laval Research Centre, Quebec City, QC, Canada, for the protection of human subjects and was conducted in accordance with the Helsinki Declaration of 1975.

Plasmids

SIN retroviral vectors were constructed based on the MFG plasmid.38 Briefly, the viral enhancers (379 bp) located in the 3’ LTR region were deleted by NheI/Sacl digestion. The digested vector was then blunt-ended with T4 DNA polymerase and ligated. The 5’ LTR was modified by replacing the HindIII/Sacl U3 region by different enhancer-promoter regions: the CMV (509 bp), the RSV (232 bp), and the CAG (644 bp).39 The human EF-1z, short version (245 bp), was then cloned in the three vectors as an internal promoter in XhoI/BamHI. In each vector, the GFP cDNA construct (720 bp) was inserted downstream of the EF-1z promoter in BamHI to generate SIN.CMV, SIN.RSV, and SIN.CAG. Three other versions of the SIN.CMV were also

study, SIN retroviral plasmids were stably introduced in the Phoenix packaging cells with the piggybac transposon system. Viral titers obtained with this method were around \(10^6\) IU/mL and were comparable to the titer of a non-SIN vector produced in PG13 cells. It was not clear whether the piggybac transposon system was superior to the standard transfection method, as there was no such comparison in the study. A2 Titeres around \(10^7\) IU/mL were achieved with our SIN GFP and PIGA vectors in the 293Vec cells, which are comparable to titers obtained with non-SIN vectors in the same packaging cells.31–34 Then, we assume that 10-fold higher titers were achieved with SIN vectors produced from 293Vec cells, as they release at least 10 times more virus than PG13 cells.34 In the second study, the SIN vector was introduced in the packaging cells by recombinase-mediated cassette exchange in a predefined locus of elevated transcription activity. With this strategy, SIN vector titers of \(2 \times 10^7\) IU/mL and \(4 \times 10^6\) IU/mL were obtained with a T cell receptor and a COL7A1 cDNA transgene, respectively. These titers are very similar to those found in our study. The advantage of this strategy is that only one copy of the vector is present in the packaging cell, which would facilitate its characterization. However, this strategy is cumbersome, as two cloning steps were necessary to obtain the final vector producer cells. The first cloning was for the exchange of the vector, and the second one was to remove the residual sequences that were required for targeting and selection.57 The 293Vec cells were also used in this study, suggesting that the potency of this packaging cell line to release high-titer vectors is key for the production of SIN vectors.

FIGURE 4. Effect of WPRE in SIN COL7A1 and PIGA Vectors Produced in Transient Transfections

Titers of COL7A1 vectors were assessed by measuring collagen VII expression by FACS analysis of HT-1080-transduced cells. Titers of PIGA vectors were assessed by measuring CD59 expression by FACS analysis of K562(PIG-)-transduced cells. Values presented are the mean ± SD of three independent experiments. **p < 0.01.
The pMD2iHygror plasmid was constructed to select stable retrovirus cations or have been synthesized by GenScript. PMD2iZeor plasmid digested by with media supplemented with 10% fetal calf serum (FCS) (Life Technologies from one RDEB patient as described elsewhere.43 isolated from a punch biopsy by the two-step thermolysine and sequences. All cell lines and primary cells.

SIN therapeutic retroviral vectors were also constructed with PIGA and COL7A1 optimized cDNA sequences (GenScript, Township, NJ, USA). The cDNA fragments were introduced in BamHI in place of the WPRE sequence (611 bp) was cloned downstream of the 3′ LTR in SmaI. A BGH poly(A) fragment (256 bp) was then cloned downstream of the 3′ LTR in AfIII to also improve the termination efficacy of the genomic RNA.

A cassette containing a BSD′ gene under the control of the HSV-TK promoter was introduced in the SIN.CMV.W.SV-BGH plasmid to facilitate the generation of stable retrovirus packaging cell clones.

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Table 1. Comparison between Hygromycin and Blasticidin Selection Systems in 293Vec-RD114-Transfected Packaging Cells

| Experiment | PCN | GFP Mean Fluorescence | Titer (10^8 IU/mL) | PCN | GFP Mean Fluorescence | Titer (10^8 IU/mL) |
|------------|-----|------------------------|--------------------|-----|------------------------|--------------------|
| 1          | 2.5 | 16.8                   | 3.4                | 13.7| 36.8                   | 8.1                |
| 2          | 2.4 | 14.2                   | 3.7                | 7   | 32.3                   | 7.7                |
| 3          | 3.9 | 16.7                   | 3.5                | 9.1 | 31.8                   | 9.2                |

PCN, plasmid copy number.

constructed. First, the WPRE sequence (661 bp) was cloned downstream of GFP to generate SIN.CMV.W. To further improve the transcriptional termination, the SIN.CMV.W.SV vector was constructed by inserting a SV40 poly(A) blunt fragment (222 bp) in the R sequence of the 3′ LTR in SmaI. A BGH poly(A) fragment (256 bp) was then cloned downstream of the 3′ LTR in AfIII to also improve the termination efficacy of the genomic RNA.

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SIN therapeutic retroviral vectors were also constructed with PIGA and COL7A1 optimized cDNA sequences (GenScript, Township, NJ, USA). The cDNA fragments were introduced in BamHI in place of the WPRE and the two poly(A) sequences. Versions without WPRE were also constructed similarly.

Fragments used in the constructions of all SIN vectors have been isolated from existing plasmids by restriction digests or PCR amplifications or have been synthesized by GenScript.

The pMD2iHygro plasmid was constructed to select stable retrovirus producer cells by co-transfection with SIN GFP vectors. A Sall/XbaI iHygro cassette was cloned instead of the iZeo cassette in the PMD2iZeo plasmid digested by Xhol/XbaI. The iHygro fragment is a Hygro gene linked to an encephalomyocarditis virus internal ribosomal entry site.

For the inactivation of the PIGA gene, the PX330-Pig plasmid was constructed by inserting a guide pair of the following annealed oligonucleotides: 5′-caccgctcagtgcctgattgaaaga-3′ and 5′-aaactctttcaatcaggcatttccttgattagagat-3′. This plasmid co-expresses the S. pyogenes Cas9 nuclease along with the PIGA guide RNA.

Cell Lines and Primary Cells

Human RDEB keratinocytes and fibroblasts used in this study were isolated from a punch biopsy by the two-step thermolysine and trypsin method from one RDEB patient as described elsewhere.43

All cell lines and primary fibroblasts used in this study were cultured with media supplemented with 10% fetal calf serum (FCS) (Life Technologies, Grand Island, NY, USA) and antibiotics. Human primary fibroblasts, HT-1080 cells (ATCC, CCL-121), 293T cells (ATCC, CRL-11268), 293Vec-RD114 cells, 293Vec-Ampho cells, and their derivatives containing the GFP, the COL7A1, and the PIGA SIN vectors were cultured with DMEM (Wisent, Saint-Jean-Baptiste, QC, Canada). Human peripheral blood lymphocytes, K562 cells, and K562(PIG-) cells were cultured with RPMI (Wisent). Human primary keratinocytes were cultured with DMEM/Ham’s F12 (3:1) (Life Technologies) supplemented with 5% FetalClone II Serum (Hyclone, Logan, UT, USA), insulin (5 μg/mL; Sigma, St. Louis, MO, USA), hydrocortisone (0.4 μg/mL; Calbiochem, San Diego, CA, USA), isoproterenol hydrochloride (0.212 μg/mL; Sandoz Canada, Boucherville, QC, Canada), epidermal growth factor (10 ng/mL; Austral Biologicals, San Ramon, CA, USA), penicillin G (100 IU/mL), and gentamicin (25 μg/mL; Sigma). For all experiments, keratinocytes were co-cultured with irradiated human fibroblast feeder layers as previously described.43

Transfections and Virus Productions

The K562(PIG-) cell line was established by electroporating K562 cells with the PX-330-Pig plasmid using a Gene Pulser electroporation system (Bio-Rad Laboratories, Hercules, CA, USA).72 A few days after recovery, cells were stained with a CD59 antibody labeled fluorescein isothiocyanate (FITC; Cederlane, Burlington, ON, Canada), and CD59-negative cells were selected by fluorescence-activated cell sorting (FACS) with the BD FACSaria II (BD Biosciences, San Jose, CA, USA). One week later, a second enrichment by FACS was performed, followed by cloning in 96-well plates. After a 3-week culture, the clone K562(PIG-) was selected and used for subsequent experiments.

Recombinant retroviruses were generated by transient transfection of 293T cells in 6-well plates using the polyethyleneimine transfection procedure. A total amount of 2 μg plasmid DNA containing 0.5 μg pMD2G/Zeopig (gag/pol expression plasmid), 0.5 μg pMD2.G (VSV-G envelope expression plasmid), and 1 μg transfer vector was transfected. Sixteen hours after transfection, the medium was replaced by 1 mL DMEM. On the next day, the culture supernatant was harvested, aliquoted, and stored at −80°C for later use.

Stable retrovirus producer cells were generated by transfection using the calcium phosphate procedure. Subconfluent 293Vec-RD114 or 293Vec-Ampho cells plated in a 10-cm dish were transfected with 20 μg BSD′ SIN vector plasmids. Two days later, cells were selected in media supplemented with blasticidin (8 μg/mL; Invitrogen, moleculartherap.jpg
For the production of virus from stable producer cells, 5/C2 was transfected with SIN GFP plasmids to transfect 293Vec-RD114 cells. Two days later, cells were selected in media supplemented with SIN GFP plasmids from a healthy donor were purified with Ficoll-Hypaque sedimentation of peripheral blood lymphocytes. Cells were then incubated for 2 h in medium with 10% FCS in a 10-cm dish, and non-adherent cells were then harvested and activated by beads coated with anti-CD3 and anti-CD28 antibodies for 48 h (Invitrogen) at a cell/bead ratio of 1/3 in medium supplemented with interleukin-2 (30 U/mL; Peprotech, Montreal, QC, Canada). Activated lymphocytes were then added onto retronectin-coated 96-well plates at 2.5 × 10⁴ cells per well with virus at a MOI of 5.

Transduction of Human Primary T Lymphocytes and RDEB Human Keratinocytes and Fibroblasts

Purification and transduction of human primary T lymphocytes were performed as previously described. Briefly, T cells isolated from a healthy donor were purified with Ficoll-Hypaque sedimentation of peripheral blood lymphocytes. Cells were then incubated for 2 h in medium with 10% FCS and non-adherent cells were then harvested and activated by beads coated with anti-CD3 and anti-CD28 antibodies for 48 h (Invitrogen) at a cell/bead ratio of 1/3 in medium supplemented with interleukin-2 (30 U/mL; Peprotech, Montreal, QC, Canada). Activated lymphocytes were then added onto retronectin-coated 96-well plates at 2.5 × 10⁴ cells per well with virus at a MOI of 5.

RDEB human fibroblasts and keratinocytes null for collagen type VII were transduced with the SIN COL7A1 vector. Cells were plated at a density of 13,000 cells per square centimeter in 12-well plates, as previously described. Briefly, fibroblasts were incubated for 2 h, and keratinocytes were incubated for 4 h at 37°C at 8% CO₂. Then, fibroblasts and keratinocytes were transduced at a MOI of 1 in the presence of 10 µg/mL of the EF-C peptide (GenScript), a better transduction enhancer than PB. Primary cell transduction efficiency was evaluated by FACS analysis, as described earlier in the section titled Titrations.

Real-Time qPCR

Genomic DNA (gDNA) was extracted using the DNeasy Tissue Kit (QIAGEN, Valencia, CA, USA). Real-time qPCR was performed using a Platinum SYBR Green qPCR SuperMix UDG kit (Thermo Fisher Scientific, Waltham, MA, USA) with a LightCycler 480 (Roche, Laval, QC, Canada). The plasmid integrated copy number was determined from a standard curve generated with the linearized plasmid SinVec.GFP, using primers specific for the WPRE sequence, or with infection, p is the percentage of fluorescent-positive cells determined by flow cytometry, V is the viral volume applied, and D is the virus dilution factor. Titers were calculated when the percentage of fluorescent-positive cells constituted between 2% to 20%.

Titrations

For all vectors, titers were determined by scoring fluorescent-positive target cells by FACS analysis. For the GFP and COL7A1 vectors, HT-1080 cells were inoculated at a density of 10⁵ cells per well in 24-well plates. On the next day, the medium from each well was replaced with 1 mL serial dilutions of virus supernatants containing polybrene (PB; 8 µg/mL). Forty-eight to 72 h later, cells transduced with GFP vectors were trypsinized and analyzed for fluorescence by FACS. Cells transduced with the COL7A1 vectors were trypsinized and permeabilized with Perm/Wash Buffer (BD Biosciences). Cells were incubated with a mouse monoclonal antibody raised against human collagen VII (LH7.2; 1:1,000, Sigma), followed by a phycoerytrin antibody (1:1,000; BioLegend), and the fluorescence was analyzed by FACS. For the titration of the PIGA vectors, 100,000 K562(PIG-) cells per well in 48-well plates were transduced with serial dilutions of vectors in the presence of PB. Two days later, cells were stained with a FITC-labeled CD59 antibody (1:1,000; Cedarlane).

Vector titers were calculated using the following formula (N × P) × 2/(V × D), in which N is the cell number at the day of production of SIN Vectors from Stable Packaging Cell Clones Selected in Blasticidin

(A) Screening titers of vectors produced from 293Vec-RD114 clones transfected with SinVec.GFP and SinVec.PIGA and from 293Vec-Ampho clones transfected with SinVec.COL7A (-W). (B) Titrations of SinVec.GFP and SinVec.PIGA vectors produced from the best 293Vec-RD114 clones and titration of the SinVec.COL7A1 (-W) produced from the best 293Vec-Ampho clone.
Figure 6. Transduction of Human Lymphocytes, RDEB Keratinocytes and Fibroblasts with SIN Vectors
(A and B) FACS analysis of primary cells was measured 3 days after transduction (A) of lymphocytes with the SinVec.GFP vector at a MOI of 5 and (B) of RDEB keratinocytes and fibroblasts with the SinVec.COL7A1 (-W) vector at a MOI of 1.

CONFLICTS OF INTEREST
M.C. and K.G. are inventors of a patent on the 293Vec technology and are shareholders of Biovec Pharma. The remaining authors declare no competing interests.

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SUPPLEMENTAL INFORMATION
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AUTHOR CONTRIBUTIONS
K.G. and M.C. designed the experiments. K.G., M.B.-W., S.R., A.D.-P., M.B., and M.C. generated reagents, performed the experiments, and analyzed data. M.C. wrote the manuscript. All authors revised the manuscript.
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