Generate DNA fragments

Synthesize a concatemer array

Introduce the concatemer array into GPRG cells

Select for transfected clones

Perform the single colony isolation

Evaluate viral production of each cell clone

FIG. 2

(57) Abrégé/Abstract:
The present disclosure provides a method of generating a stable producer cell line. The generation of stable producer cell lines, such as those provided in accordance with the present invention, increases the reproducibility and ease of creating high titer lentiviral stocks while easing biosafety concerns and the variation in expressed envelope proteins defines the tropism of the generated virus. The present disclosure also provides for a novel lentiviral transfer vector plasmid.
ABSTRACT

The present disclosure provides a method of generating a stable producer cell line. The generation of stable producer cell lines, such as those provided in accordance with the present invention, increases the reproducibility and ease of creating high titer lentiviral stocks while easing biosafety concerns and the variation in expressed envelope proteins defines the tropism of the generated virus. The present disclosure also provides for a novel lentiviral transfer vector plasmid.
BIO-PRODUCTION OF LENTIVIRAL VECTORS

CROSS-REFERENCE TO RELATED APPLICATIONS

[0001] The present application claims the benefit of the filing date of United States Provisional Patent Application No. 62/161,133 filed May 13, 2015; and also claims the benefit of the filing date of United States Provisional Patent Application No. 62/161,152, filed May 13, 2015, the disclosures of which are hereby incorporated by reference herein in their entireties.

FIELD OF DISCLOSURE

[0002] This disclosure generally relates to the fields of molecular biology and virology. In particular, the disclosure relates to the bio-production of lentiviral vectors and lentiviral transfer plasmids.

STATEMENT OF INDUSTRIAL APPLICABILITY

[0003] The present disclosure has industrial applicability in the field of gene therapeutics and bio-manufacturing.

BACKGROUND OF THE DISCLOSURE

[0004] HIV-1 is the causative agent of Acquired Immunodeficiency Syndrome (AIDS) with of the order of 30 million individuals infected world-wide. HIV causes the immune system to fail and increases the probability of death due to opportunistic infections. HIV infection is a major global health problem as evidenced by its designation as a pandemic by the World Health Organization. Most people who are infected with HIV, particularly in the developing world, eventually develop AIDS, which claims the lives of more than one million people every year.

[0005] HIV-1 belongs to the retroviridae family of viruses, and is an enveloped virus whose genome consists of two single stranded RNA molecules (ssRNA). The primary target of HIV-1 is CD4+ expressing cells, such as CD4+ T cells. A glycoprotein of the HIV-1 virus interacts with the CD4 molecule of target cells and with chemokine co- receptors, CCRS or CXCR4 on the surface of target cells. Following fusion and entry into the target cell, the nucleocapsid containing the viral
genome dissociates, releasing the contents of the virus, including the ssRNA, into the cytoplasm. A reverse transcriptase (RT) enzyme of HIV-1 synthesizes viral double stranded DNA (dsDNA) from the ssRNA genome. Following synthesis of the double stranded HIV-1 DNA molecule, the HIV-1 DNA is integrated into the host genome.

[0006] The integrated HIV-1 DNA is flanked by identical 5' and 3' long terminal repeat sequences (LTR) from which HIV-1 can initiate transcription of the integrated HIV-1 genome. Transcription of the viral DNA requires transcription factors, such as NF-kB, which are upregulated in activated T cells. As a consequence, viral transcription is most active in the T cell following activation of the T cell, such as during infection. Viral RNA resulting from transcription of the integrated HIV-1 genome is subsequently translated and packaged into virus particles which then exit the cell to become infectious virus.

[0007] Therapy for HIV-1 infection includes combination antiretroviral therapy (cART). cART, which includes combinations of nucleoside analogue reverse transcriptase inhibitors, protease inhibitors, non-nucleoside reverse transcriptase inhibitors, integrase and fusion inhibitors, slows HIV progression. This, in turn, dramatically decreases the morbidity and mortality rate from HIV/AIDS in regions of the world where the therapy is available. However, cART does not cure or completely eliminate all the symptoms of HIV/AIDS. Also, cART therapy can be compromised by drug resistant mutations, and has a range of side effects which can be serious and which appear to be cumulative. Further, interruption of cART therapy almost invariably leads to the re-emergence of detectable viral replication and the progression to AIDS and has been shown to be associated with an increased incidence of all causes of mortality and serious non AIDS events. For these reasons, as well as the high cost of cART and need for strict adherence, such therapy can be relatively ineffective for a large number of patients.

[0008] HIV-based lentiviral vectors are rapidly becoming the retrovirus vector system of choice for research and clinical gene transfer applications. The enhanced ability of lentiviral vectors to transduce both quiescent stem cells and non-dividing terminally differentiated cells has led to the development of a wide range of therapeutic gene delivery vectors, as well as promising research tools, such as short hairpin RNA
(shRNA) gene knockdown libraries and vectors for induction of pluripotency in terminally differentiated cells. Early gamma-retroviral clinical gene therapy vectors restored immune function in patients with X-linked severe combined immunodeficiency (SCID-X1), but they were subsequently found to cause proliferative disorders via transactivation of proto-oncogenes. Newer lentiviral vector designs may significantly reduce that risk, and they await clinical testing for final validation of their predicted safety. The field remains in flux and the outcomes of the clinical testing are unpredictable.

[0009] Producing SIN - lentiviral vectors at scales to support clinical trials is an important challenge within the field. While gamma-retroviral vectors can be produced by either transient transfection or the generation of stable producer cell lines, lentiviruses require the expression of multiple cytotoxic accessory genes, which makes the generation of producer cells more complicated (Greene et al., Transduction of Human CD34+ Repopulating Cells with a Self – Inactivating Lentiviral Vector for SCID-X1 Produced at Clinical Scale by a Stable Cell Line, HGTM, 23, 297-308 (October 2012), which is hereby incorporated by reference in its entirety). Transient transfection is instead the current technology for pilot production of LV, which is impractical for very large-scale applications under a safety, cost, and reproducibility standpoint. In fact, this technology is expensive, is difficult to standardize and scale-up, and suffers from batch-to-batch variability and low reverse transcriptase fidelity (Storaiuolo et al., RD2-MolPack-Chim3, a Packaging Cell Line for Stable Production of Lentiviral Vectors for Anti-HIV Gene Therapy, HGTM, 24:228-240 (August 2013), which is hereby incorporated by reference in its entirety).

SUMMARY OF THE DISCLOSURE

[0010] In one aspect of the present disclosure is a plasmid comprising a nucleotide sequence having at least 90% identity to that of SEQ ID NO: 1. In some embodiments, the nucleotide sequence has at least 95% identity to that of sequence of SEQ ID NO: 1. In some embodiments, a lentiviral vector is derived from the plasmid. In some embodiments, the derived lentiviral vector comprises one or more additional sequences encoding a short hairpin RNA for down-regulation of an HIV-1 co-receptor
and/or an HIV-1 fusion inhibitor. In some embodiments, the lentiviral vector derived therefrom is LVsh5/C46 (as defined herein).

[0011] In another aspect of the present disclosure is a plasmid comprising the nucleotide sequence of SEQ ID NO: 1. In some embodiments, a lentiviral vector is derived from the plasmid. In some embodiments, the derived lentiviral vector comprises one or more additional sequences encoding a short hairpin RNA for down-regulation of an HIV-1 co-receptor (e.g. CCR5) and/or an HIV-1 fusion inhibitor (e.g. C46). Information regarding CCR5 and C46, including their nucleotide sequences, is set forth further in US Patent Publication No. US2012/0201794, the disclosure of which is incorporated by reference herein in its entirety.

[0012] In another aspect of the present disclosure is a plasmid comprising a nucleotide sequence having at least 80% identity to that of SEQ NO: 2. In some embodiments, the plasmid comprises a nucleotide sequence having at least 90% identity to that of SEQ NO: 2. In some embodiments, the plasmid comprises a nucleotide sequence having at least 95% identity to that of SEQ NO: 2. In some embodiments, the plasmid comprises a nucleotide sequence having at least 97% identity to that of SEQ NO: 2. In some embodiments, the plasmid comprises the nucleotide sequence of SEQ NO: 2. In some embodiments, a lentiviral vector is derived from the plasmid. In some embodiments, the derived lentiviral vector comprises one or more additional sequences encoding a short hairpin RNA for down-regulation of an HIV-1 co-receptor (e.g. CCR5) and/or an HIV-1 fusion inhibitor (e.g. C46).

[0013] In another aspect of the present disclosure is a plasmid having a sequence that differs by not more than 500 nucleotides from the sequence set forth in SEQ ID NO: 1 (e.g. non-consecutive or consecutive). In another aspect of the present disclosure is a plasmid having a sequence that differs by not more than 250 nucleotides from the sequence set forth in SEQ ID NO: 1 (e.g. non-consecutive or consecutive). In another aspect of the present disclosure is a plasmid having a sequence that differs by not more than 150 nucleotides from the sequence set forth in SEQ ID NO: 1 (e.g. non-consecutive or consecutive). In another aspect of the present disclosure is a plasmid having a sequence that differs by not more than 100 nucleotides from the sequence set
forth in SEQ ID NO: 1 (e.g. non-consecutive or consecutive). In some embodiments, the sequence differs by not more than 50 nucleotides from the sequence set forth in SEQ ID NO: 1 (e.g. non-consecutive or consecutive).

[0014] In another aspect of the present disclosure is a plasmid comprising between about 6500 nucleotides and about 6750 nucleotides, and wherein the plasmid comprises a sequence or fragment thereof having at least 90% identity to that of SEQ ID NO: 2. In some embodiments, the plasmid comprises between about 6600 nucleotides and about 6700 nucleotides. In some embodiments, the plasmid comprises about 6611 nucleotides.

[0015] In another aspect of the present disclosure is a plasmid as set forth in FIG. 11 as pUC57-TL20c. In some embodiments, a lentiviral vector is derived from the plasmid. In some embodiments, the derived lentiviral vector comprises one or more additional sequences encoding a short hairpin RNA for down-regulation of an HIV-1 co-receptor and/or an HIV-1 fusion inhibitor.

[0016] In another aspect of the present disclosure is a plasmid comprising a multiple cloning site consisting essentially of the BstBI, MluI, NotI, and ClaI restriction endonuclease sites. In some embodiments, the plasmid further comprises a nucleotide sequence encoding a packaging signal; a nucleotide sequence encoding a central polyuridine tract; a nucleotide sequence encoding a Rev response element; and a nucleotide sequence encoding a self-inactivating long terminal repeat. In other embodiments, the plasmid comprises a multiple cloning site consisting of the BstBI, MluI, NotI, and ClaI restriction endonuclease sites. In some embodiments, a lentiviral vector is derived from the plasmid. In some embodiments, the derived lentiviral vector comprises one or more additional sequences encoding a short hairpin RNA for down-regulation of an HIV-1 co-receptor and/or an HIV-1 fusion inhibitor.

[0017] In another aspect of the present disclosure is a plasmid comprising (a) a nucleotide sequence encoding a packaging signal; (b) a nucleotide sequence encoding a central polyuridine tract (cPPT); (c) a nucleotide sequence encoding a Rev response element; (d) a nucleotide sequence encoding a self-inactivating long terminal repeat; and (e) a nucleotide sequence encoding a multiple cloning site having restriction sites for the enzymes BstBI, Mlu I, Not I, and Cla I. In some embodiments, the nucleotide
sequence encoding the packing signal comprises the sequence of SEQ ID NO: 3. In some embodiments, the nucleotide sequence encoding the central polypurine tract (cPPT) comprises the sequence of SEQ ID NO: 4. In some embodiments, the nucleotide sequence encoding the Rev response element comprises the sequence of SEQ ID NO: 5. In some embodiments, the nucleotide sequence encoding the self-inactivating long terminal repeat comprises the sequence of SEQ ID NO: 6. In some embodiments, the nucleotide sequence encoding the multiple cloning site comprises the sequence of SEQ ID NO: 7.

[0018] In another aspect of the present disclosure is a plasmid comprising (a) a packaging sequence, the packaging sequence present from about nucleotide 762 to about nucleotide 1104 of a nucleotide sequence of the plasmid; (b) a central polypurine tract, the central polypurine tract present from about nucleotide 1121 to about nucleotide 1597 of the plasmid nucleotide sequence; (c) a Rev response element, the Rev response element present from about nucleotide 1598 to about nucleotide 2366 of the plasmid nucleotide sequence; (d) a self-inactivating long terminal repeat, the self-inactivating long terminal repeat present from about nucleotide 409 to about nucleotide 589 of the plasmid nucleotide sequence; and (e) a multiple cloning site, the multiple cloning site present from about nucleotide 2376 to about nucleotide 2400 of the plasmid nucleotide sequence. In some embodiments, the plasmid nucleotide sequence comprises a sequence having at least 90% identity to that of SEQ ID NO: 1. In some embodiments, a nucleotide sequence encoding the packing signal comprises the sequence of SEQ ID NO: 3. In some embodiments, nucleotide sequence encoding the central polypurine tract (cPPT) comprises the sequence of SEQ ID NO: 4. In some embodiments, a nucleotide sequence encoding the Rev response element comprises the sequence of SEQ ID NO: 5. In some embodiments, a nucleotide sequence encoding the self-inactivating long terminal repeat comprises the sequence of SEQ ID NO: 6. In some embodiments, a nucleotide sequence encoding the multiple cloning site comprises the sequence of SEQ ID NO: 7.

[0019] In another aspect of the present disclosure is a plasmid comprising a multiple cloning site comprising the BstBI, MluI, NotI, and ClaI restriction endonuclease sites, and wherein the plasmid comprises a nucleotide sequence having at
least 80% identify to that of SEQ ID NO: 1. In some embodiments, a lentiviral vector is derived from the plasmid. In some embodiments, the derived lentiviral vector comprises one or more additional sequences encoding a short hairpin RNA for down-regulation of an HIV-1 co-receptor and/or an HIV-1 fusion inhibitor.

[0020] In another aspect of the present disclosure is a plasmid comprising a nucleotide sequence encoding a vector backbone having at least 95% identity to that of SEQ ID NO: 2, and wherein the vector backbone is flanked by at least two additional restriction endonuclease sites, the at least two additional restriction endonuclease sites independently are selected from the group consisting of SfiI and Bsu36I. In some embodiments, a lentiviral vector is derived from the plasmid. In some embodiments, the derived lentiviral vector comprises one or more additional sequences encoding a short hairpin RNA for down-regulation of an HIV-1 co-receptor and/or an HIV-1 fusion inhibitor.

[0021] In another aspect of the present disclosure is plasmid comprising a nucleotide sequence encoding a vector backbone having at least 90% identity to that of SEQ ID NO: 2, the vector backbone comprising a multiple cloning site having BstBI, MluI, NotI, and ClaI restriction endonuclease sites, wherein the plasmid further comprises a tetracycline repressible promoter upstream of the vector backbone. In yet another aspect of the present disclosure is plasmid comprising a nucleotide sequence encoding a vector backbone having at least 85% identity to that of SEQ ID NO: 2, the vector backbone consisting of a multiple cloning site having BstBI, MluI, NotI, and ClaI restriction endonuclease sites,

[0022] In another aspect of the present disclosure is a cell comprising a plasmid or a lentiviral vector derived therefrom as described herein. In some embodiments, the cell is a hematopoietic progenitor/stem cell, a monocyte, a macrophage, a peripheral blood mononuclear cell, a CD4+ T lymphocyte, a CD8+ T lymphocyte, or a dendritic cell. In another aspect of the present invention is a kit comprising hematopoietic progenitor/stem cells in a first container and the plasmid of FIG. 11 or a lentiviral vector derived therefrom.

[0023] In another aspect of the present disclosure is a method of producing a stable producer cell line comprising: (a) synthesizing a lentiviral vector by cloning one
or more genes into a plasmid as described herein, e.g. pUC57-TL20c; (b) generating DNA fragments from the synthesized lentiviral vector; (c) forming a concatemeric array from the generated DNA fragments of the synthesized lentiviral vector and from DNA fragments from an antibiotic resistance cassette plasmid; (d) transfecting a GPR, GPRG, GPRT, GPRGT or GPRT-G packing cell line or a derivative thereof with the formed concatemeric array; and (e) isolating one or more stable producer cell line clones. In some embodiments, the method further comprises inducing the stable producer cell line to produce the lentiviral vector.

[0024] In another aspect of the present disclosure is a method of producing a stable producer cell line comprising: (a) synthesizing a lentiviral vector which encodes a short hairpin RNA for down-regulation of an HIV-1 co-receptor and which encodes an HIV-1 fusion inhibitor, the lentiviral vector synthesized by cloning cDNA encoding both the short hairpin RNA and the fusion inhibitor into a plasmid as described herein; (b) generating DNA fragments from the synthesized lentiviral vector; (c) forming a concatemeric array from the generated DNA fragments from the synthesized lentiviral vector and from DNA fragments from an antibiotic resistance cassette plasmid; (d) transfecting a GPR, GPRG, GPRT, GPRGT or GPRT-G packing cell line or a derivative thereof with the formed concatemeric array; and (e) isolating one or more stable producer cell line clones. In some embodiments, the method further comprises inducing the stable producer cell line to produce the lentiviral vector which encodes a short hairpin RNA for down-regulation of an HIV-1 co-receptor and which encodes an HIV-1 fusion inhibitor (LVsh5/C46).

[0025] In another aspect of the present disclosure is a method of harvesting vector supernatant from a stable producer cell line, wherein the vector supernatant is harvested about every 48 hours. In another aspect of the present disclosure is a method of harvesting vector supernatant from a stable producer cell line, wherein the vector supernatant is harvested every 40 to 56 hours.

[0026] In another aspect of the present disclosure is a method of harvesting vector supernatant comprising the LVsh5/C46 lentiviral vector, wherein the vector supernatant is harvest about every 48 hours.

[0027] In another aspect of the present disclosure is a stable producer cell line suitable for producing LVsh5/C46. In some embodiments, the stable producer cell line
is based on the GPRG packaging cell line. In some embodiments, the stable producer cell line based on the GPRT packaging cell line. In some embodiments, the stable producer cell line based on the GPRT-G packaging cell line.

[0028] In another aspect of the present disclosure is a stable producer cell line suitable for producing a self-inactivating lentiviral vector having at least 90% identity to that of SEQ ID NO: 8. In some embodiments, the stable producer cell line is based on the GPRG packaging cell line. In some embodiments, the stable producer cell line based on the GPRT packaging cell line. In some embodiments, the stable producer cell line based on the GPRT-G packaging cell line. In some embodiments, the stable producer cell line based on the GPRT-G packaging cell line.

[0029] In another aspect of the present disclosure is a concatemeric array comprising DNA fragments from a first plasmid and a second plasmid; the first plasmid derived from pUC57-TL20c; the second plasmid comprising a bleomycin antibiotic resistance cassette; wherein a ratio of the DNA fragments from the first plasmid to the second plasmid ranges from about 50:1 to about 1:50. In some embodiments, the ratio of the DNA fragments from the first plasmid to the second plasmid ranges from about 25:1 to about 1:25. In some embodiments, the ratio of the DNA fragments from the first plasmid to the second plasmid ranges from about 15:1 to about 1:15.

[0030] In another aspect of the present disclosure is a stable producer cell line, the stable producer cell line generated by transfecting a packaging cell line selected from the group consisting of GPR, GPRG, GPRT, GPRT-G and a derivative thereof with a concatemeric array, the concatemeric array comprising DNA fragments from a first plasmid and a second plasmid; the first plasmid derived from pUC57-TL20c; the second plasmid comprising a bleomycin antibiotic resistance cassette; wherein a ratio of the DNA fragments from the first plasmid to the second plasmid ranges from about 25:1 to about 1:25. In some embodiments, the stable producer cell line produces LVsh5/C46. In some embodiments, the LVsh5/C46 is capable of being harvested about every 48 hours.
[0031] In another aspect of the present disclosure is an isolated vector having the plasmid map of FIG. 11 which comprises a multiple cloning site that comprises the nucleotide sequence set forth in SEQ ID NO: 7.

[0032] In another aspect of the present disclosure is a kit comprising (1) a plasmid as described herein, and (2) a bleomycin resistance (ble) cassette. In some embodiments, the kit further comprises instructions for preparing a lentiviral vector and/or a concatemeric array, such as according to the procedures described herein.

[0033] In another aspect of the present disclosure is a kit comprising (a) a lentiviral transfer vector plasmid as described herein; and (b) a packaging cells. In some embodiments, the packaging cells are selected from the group consisting of GPR, GPRG, GPRT, GPRTG, and a derivative thereof. In some embodiments, the kit further comprises a bleomycin resistance (ble) cassette. In some embodiments, the kit further comprises instructions for preparing a lentiviral vector and/or a concatemeric array, such as according to the procedures described herein.

[0034] In another aspect of the present disclosure is a lentiviral vector derived from a plasmid as described herein. In some embodiments, the lentiviral vector comprises at least one additional nucleotide sequence. In some embodiments, the at least one additional nucleotide sequence is selected from the group consisting of a nucleotide sequence which encodes a short hairpin RNA for down-regulation of an HIV-1 co-receptor and a nucleotide sequence which encodes an HIV-1 fusion inhibitor. In some embodiments, the lentiviral vector is LVsh5/C46. In some embodiments, the lentiviral vector comprises a sequence having at least 95% identity to that of SEQ ID NO: 8.

[0035] In another aspect of the present disclosure is a pharmaceutical composition comprising the lentiviral vector described above and a pharmaceutically acceptable carrier. In some embodiments, the pharmaceutically acceptable carrier includes solvents, buffers, solutions, dispersion media, coatings, antibacterial and antifungal agents, isotonic and absorption delaying agents and the like acceptable for use in formulating pharmaceuticals, such as pharmaceuticals suitable for administration to humans. Methods for the formulation of compounds with pharmaceutical carriers are known in the art and are described in, for example, in Remington's Pharmaceutical Science, (17th ed. Mack Publishing Company, Easton, Pa. 1985); and Goodman &
Gillman's: The Pharmacological Basis of Therapeutics (11th Edition, McGraw-Hill Professional, 2005); the disclosures of each of which are hereby incorporated herein by reference in their entirety.

**BRIEF DESCRIPTION OF THE DRAWINGS**

[0036] Figure 1 illustrates that the same lentiviral vector was produced repeatedly by either transient transfection on HEK293T/17 cells according to established procedures, or using a GPRG-based stable producer cell line. Vector containing media (VCM) was concentrated 100x by ultracentrifugation and lentiviral (LV) titer was determined by gene transduction assay.

[0037] Figure 2 is a flowchart illustrating a method of generating a stable producer cell line and for harvesting lentiviral vectors produced from the generated stable producer cell line.

[0038] Figure 3 illustrates the assessment of producer cell line stability for two different cell line MWCBs over a three-month period of continuous passage. At regular intervals, LV was induced by tetracycline (TET) removal and VCM was assessed for LV titer by gene transduction assay. Both cell lines were stable and able to produce LV in excess of 1e6/ml over the three-month period and for in excess of about 25 passages.

[0039] Figures 4 illustrates the kinetics of lentiviral vector production following induction by removal of TET. Vector titer was assessed in VCM by gene transduction assay. In all instances, GPRG-based stable producer cell lines have been able to maintain LV production at levels above 1e6 TU/ml (unconcentrated) for at least 5 days following induction.

[0040] Figure 5 illustrates the kinetics of lentiviral vector production from stable cell lines. (A) During vector production, the medium was replaced with a fresh medium on a daily basis (■) or every 2 days (□). (B) The total amount of LVs in the harvested medium was titrated on 293T cells. The data shown are the mean values ± SD (N = 2). TU units, transduction units.

[0041] Figure 6-(A) illustrates GPRG and 293T cells were induced in medium without doxycycline (Dox). The induced cells were stained by anti-VSVG antibodies to
detect the VSVG expression and measured by flow cytometry; (B) illustrates the ability of the GPRG to produce LV was assessed after prolonged culture.

[0042] Figure 7 illustrates lentiviral production in different culture conditions. (A) Cultured/Produced in serum-containing medium. (B, Left) Cultured in serum-containing/Produced in serum-free medium; (B, Right) Cultured/Produced in serum-free medium. D10: 500 mL DMEM/GlutaMAX™; 50 mL FBS (10% w/v); 5 mL Pen/Strep; SFM: serum-free medium.

[0043] Figure 8 sets forth a FACS analysis of 293T or TF-1a cells incubated with either fresh medium (no vector) or LVsh5/C46 vectors.

[0044] Figure 9 illustrates the quantification of lentiviral vector copy numbers in the infected cells. C46 qPCR was used to determine the vector copy number per host genome after transduction at two doses (MOI = 1 or 0.3).

[0045] Figure 10–Ghost-CCR5 cells were transduced with LVsh5/C46 vectors. The decreased level of CCR5 expression was measured by FACS.

[0046] Figure 11 sets forth a schematic diagram of pUC57-TL20.

[0047] Figure 12 illustrates an HIV-1 based lentiviral transfer vector according to some embodiments of the present disclosure. This particular transfer vector encodes a short hairpin RNA (shRNA) for down-regulation of the HIV-1 co-receptor CCR5, in combination with a HIV-1 fusion inhibitor (C46).

[0048] Figure 13 illustrates lentiviral induction from using the methods disclosed herein with and without serum. Cells cultured in serum-free media produced nearly as much virus as those cultured with 10% PBS. It is believed that the methods disclosed here may be adapted to serum-free culture environments.

[0049] Figure 14 is a flowchart illustrating a method of generating DNA fragments.

[0050] Figure 15 is a flowchart illustrating a method of synthesizing a concatemeric array.

[0051] Figure 16 is a flowchart illustrating a method of introducing a concatemeric array into a packaging cell line.
[0052] Figure 17 is a flowchart illustrating a method of selecting for transfected clones.

[0053] Figure 18 is a flowchart illustrating a method of performing a single colony isolation.

[0054] Figure 19 is a flowchart illustrating a method of evaluating viral production.

[0055] Figures 20A, 20B, and 20C, in general, describe producer cells for synthesizing TL20-Cal1-wpre and TL20-Unc-GFP vectors. Figure 20A illustrates a flow cytometry analysis of 293T cells incubated with either fresh medium (left: no vector) or TL20-Cal1-WPRE (Right) harvested from the most potent producer clone. Figure 20B illustrates a flow cytometry analysis of 293T cells incubated with either fresh medium (dark grey bar: no vector) or TL20-UbcGFP (light grey bar) harvested from the most potent producer clone. Figure 20C illustrates the distribution of measured vector titers of supernatants from the independent producer clones for making the TL20-Cal1-WPRE (left) or TL20-UbcGFP (right) vector. The vectors were titrated on 293T cells and analyzed by flow cytometry. The highest titer achieved for the vectors prepared using polyclonal producer cells (before single clonal selection) is indicated by dashed line. Legend: Ubc: Ubiquitin C promoter; GFP: enhanced green fluorescence protein.

DETAILED DESCRIPTION

[0056] In general, the present disclosure provides a method of generating a stable producer cell line. The generation of stable producer cell lines, such as those provided in accordance with the present invention, increases the reproducibility and ease of creating high titer lentiviral stocks while easing biosafety concerns and the variation in expressed envelope proteins defines the tropism of the generated virus. The present disclosure also provides for a novel lentiviral transfer vector plasmid.

[0057] As used herein, the singular terms "a," "an," and "the" include plural referents unless the context clearly indicates otherwise. Similarly, the word "or" is intended to include "and" unless the context clearly indicates otherwise.
The terms "comprising," "including," "having," and the like are used interchangeably and have the same meaning. Similarly, "comprises," "includes," "has," and the like are used interchangeably and have the same meaning. Specifically, each of the terms is defined consistent with the common United States patent law definition of "comprising" and is therefore interpreted to be an open term meaning "at least the following," and is also interpreted not to exclude additional features, limitations, aspects, etc. Thus, for example, "a device having components a, b, and c" means that the device includes at least components a, b and c. Similarly, the phrase: "a method involving steps a, b, and c" means that the method includes at least steps a, b, and c. Moreover, while the steps and processes may be outlined herein in a particular order, the skilled artisan will recognize that the ordering steps and processes may vary.

As used herein, the term "cloning" refers to the process of ligating a nucleic acid molecule into a plasmid and transferring it into an appropriate host cell for duplication during propagation of the host.

As used herein, the term "HIV" includes not only HIV-1, but also the various strains of HIV-1 (e.g. strain BaL or strain SF162) and the various subtypes of HIV-1 (e.g. subtypes A, B, C, D, F, G H, I, and K).

As used herein, the term "multiple cloning site" (MCS) refers to nucleotide sequences comprising restriction sites for the purpose of cloning nucleic acid fragments into a cloning vector plasmid. A MCS, also referred to as a poly linker or polycloning site, is a cluster of cloning sites such that many restriction enzymes are able to operate within the site. A cloning site in some embodiments is a known sequence upon which a restriction enzyme operates to linearize or cut a plasmid.

As used herein, the term "producer cell" refers to a cell which contains all the elements necessary for production of lentiviral vector particles.

As used herein, the term "packaging cell" refers to a cell which contains those elements necessary for production of infectious recombinant virus which are lacking in a recombinant viral vector or lentiviral transfer vector plasmid. Typically, such packaging cells contain one or more expression cassettes which are capable of
expressing viral structural proteins (such as gag, pol and env) but they do not contain a packaging signal.

[0064] As used herein, the terms "restriction endonuclease" or "restriction enzyme" refer to a member or members of a class of catalytic molecules that bind a cognate sequence of a nucleic acid molecule (e.g. DNA) and cleave it at a precise location within that sequence.

[0065] As used herein, the term "self-inactivating" or "SIN," used interchangeably herein, refers to a vector which is modified, wherein the modification greatly reduces the ability of the vector to mobilize once it has integrated into the genome of the recipient, thereby increasing the safety of the use of the vector as a gene delivery vector.

[0066] As used herein, the term "vector" refers to a nucleic acid molecule capable of mediating entry of, e.g., transferring, transporting, etc., another nucleic acid molecule into a cell. The transferred nucleic acid is generally linked to, e.g., inserted into, the vector nucleic acid molecule. A vector may include sequences that direct autonomous replication, or may include sequences sufficient to allow integration into host cell DNA. As will be evident to one of ordinary skill in the art, viral vectors may include various viral components in addition to nucleic acid(s) that mediate entry of the transferred nucleic acid.

[0067] **Overview of Method**

[0068] Lentiviral vectors (LVs) are important tools for gene transfer due to their efficiency and ability to stably transduce both dividing and non-dividing cells. As a result, investigators are using them as gene delivery vehicles in a wide variety of clinical applications. Nevertheless, large-scale clinical production using current good manufacturing practice (cGMP) methods comes with a set of challenges that must be considered as more clinical trials using lentiviral vectors receive regulatory approval. One important consideration in designing cGMP-compatible processes is the need to integrate regulatory considerations into manufacturing processes that are capable of producing consistent lentivirus for multiple cGMP productions. The vast majority of lentiviral vectors being used clinically has been produced by transient transfection.
Transient transfection based production is, however, often labor intensive and subject to variation. For this reason, several stable packaging cell line systems have recently been developed. While the use of these cell lines for the bio-manufacturing of LV is particularly attractive for both scalability and consistency, development of such lines is time consuming and the regulatory pathway for the cGMP use of these lines has not been firmly established.

[0069] In view of this, the present disclosure sets forth a process for the clinical production of self-inactivating lentiviral vectors (SIN-LVs). It is believed that through the use of a novel lentiviral transfer vector plasmid together with GPR, GPRG, GPRT, GPRGT or GPRT-G packaging cell lines (or a derivative or analog packaging cell line derived therefrom), that stable producer cell lines may be generated so as to enable the production of self-inactivating lentiviral vectors (e.g. LVsh5/C46). While certain embodiments and examples described herein refer to the production of LVsh5/C46, which is a self-inactivating lentiviral vector encoding a short hairpin RNA (shRNA) for down-regulation of the HIV-1 co-receptor CCR5, in combination with a HIV-1 fusion inhibitor (namely, C46), the skilled artisan will recognize that the methods described herein are suitable for the generation of stable producer cell lines capable of producing any SIN-LVs, comprising any desired or client supplied genes or sequences.

[0070] Applicants have demonstrated that compared to SIN-LVs produced by transient transfection, that the presently disclosed method (i) is capable of generating a similar quality and quantity of SIN-LVs; (ii) produces LVs that may have better potency; and (ii) maintains yields while greatly decreasing prep-to-prep variability seen with transient transfection.

[0071] **pUC57-TL.20c**

[0072] In one aspect of the present disclosure is a human immunodeficiency virus type 1 (HIV-1) based third generation, self-inactivating (SIN) lentiviral transfer vector plasmid (hereinafter referred to as "pUC57-TL.20") comprising a novel, versatile multiple cloning site (MCS) (see FIG. 11).

[0073] In some embodiments, the lentiviral vector transfer plasmid comprises a vector backbone ("TL20c") that does not itself comprise an internal promoter (hence, it
is "promoterless"). In some embodiments, the lentiviral vector transfer plasmid comprises one promoter, e.g. a tetracycline repressible promoter, upstream of the vector backbone (see FIG. 12). Without wishing to be bound by any particular theory, it is believed that the promoterless design of the vector backbone allows for the generation of a lentiviral transfer vector plasmid that enables the delivery and subsequent expression of a gene of interest from a user-determined promoter.

FIG. 11 sets forth a gene map illustrating the constituent elements of the lentiviral vector transfer plasmid. In some embodiments, the lentiviral vector transfer plasmid comprises between about 6500 nucleotides and about 6750 nucleotides. In other embodiments, the lentiviral vector transfer plasmid comprises between 6600 nucleotides and about 6700 nucleotides. In some embodiments, the vector backbone of the lentiviral transfer vector plasmid comprises between about 3850 nucleotides and about 3950 nucleotides. In some embodiments, the vector backbone of the lentiviral transfer vector plasmid comprises about 3901 nucleotides.

As shown in FIG. 11, the plasmid comprises a 5' flanking HIV LTR, a packaging signal or ψ+, a central polypurine tract (cPPT), a Rev-response element (RRE), a multiple cloning site (MCS), and a 3' flanking HIV LTR. The LTR regions further comprise a U3 and U5 region, as well as an R region.

According to certain embodiments of the disclosure, the transfer plasmid includes a self-inactivating (SIN) LTR. As is known in the art, during the retroviral life cycle, the U3 region of the 3' LTR is duplicated to form the corresponding region of the 5' LTR in the course of reverse transcription and viral DNA synthesis. Creation of a SIN LTR is achieved by inactivating the U3 region of the 3' LTR (preferably by deletion of a portion thereof, e.g. removal of a TATA sequence). The alteration is transferred to the 5' LTR after reverse transcription, thus eliminating the transcriptional unit of the LTRs in the provirus, which is believed to prevent mobilization by replication competent virus. An additional safety enhancement is provided by replacing the U3 region of the 5' LTR with a heterologous promoter to drive transcription of the viral genome during production of viral particles.

In some embodiments, the packaging signal comprises about 361 base pairs of the Gag sequence and about 448 base pairs of the Pol sequence of wild-type
HIV (e.g. HIV01 HXB2_LAI_IIIB). In some embodiments, the cPPT comprises about 85 base pairs of the Vif sequence of wild-type HIV. In some embodiments, a HIV Polypurine tract (pPu) comprises about 106 base pairs of the Nef sequence of wild-type HIV. In some embodiments, the RRE comprises about 26 base pairs of the Rev sequence, about 25 base pairs of the tat sequence, and about 769 base pairs of the Env sequence of wild-type HIV. In some embodiments, the transfer plasmid comprises a chromatin insulator and/or a beta-globulin polyadenylation signal.

[0078] In some embodiments, the nucleotide sequence encoding the packing signal comprises the sequence of SEQ ID NO: 3, or a sequence having at least 85% identity to that of SEQ ID NO: 3. In some embodiments, the nucleotide sequence encoding the packing signal comprises the sequence of SEQ ID NO: 3, or a sequence having at least 90% identity to that of SEQ ID NO: 3. In some embodiments, the nucleotide sequence encoding the packing signal comprises the sequence of SEQ ID NO: 3, or a sequence having at least 95% identity to that of SEQ ID NO: 3.

[0079] In some embodiments, the nucleotide sequence encoding the central polypurine tract (cPPT) comprises the sequence of SEQ ID NO: 4, or a sequence having at least 85% identity to that of SEQ ID NO: 4. In some embodiments, the nucleotide sequence encoding the central polypurine tract (cPPT) comprises the sequence of SEQ ID NO: 4, or a sequence having at least 90% identity to that of SEQ ID NO: 4. In some embodiments, the nucleotide sequence encoding the central polypurine tract (cPPT) comprises the sequence of SEQ ID NO: 4, or a sequence having at least 95% identity to that of SEQ ID NO: 4.

[0080] In some embodiments, the nucleotide sequence encoding the Rev response element comprises the sequence of SEQ ID NO: 5, or a sequence having at least 85% identity to that of SEQ ID NO: 5. In some embodiments, the nucleotide sequence encoding the Rev response element comprises the sequence of SEQ ID NO: 5, or a sequence having at least 90% identity to that of SEQ ID NO: 5. In some embodiments, the nucleotide sequence encoding the Rev response element comprises the sequence of SEQ ID NO: 5, or a sequence having at least 95% identity to that of SEQ ID NO: 5.
[0081] In some embodiments, the nucleotide sequence encoding the self-inactivating long terminal repeat comprises the sequence of SEQ ID NO: 6, or a sequence having at least 85% identity to that of SEQ ID NO: 6. In some embodiments, the nucleotide sequence encoding the self-inactivating long terminal repeat comprises the sequence of SEQ ID NO: 6, or a sequence having at least 90% identity to that of SEQ ID NO: 6. In some embodiments, the nucleotide sequence encoding the self-inactivating long terminal repeat comprises the sequence of SEQ ID NO: 6, or a sequence having at least 95% identity to that of SEQ ID NO: 6.

[0082] In some embodiments, the plasmid comprises a nucleotide sequence encoding a doxycycline repressible promoter that has at least 85% identity to that of SEQ ID NO: 10. In some embodiments, the plasmid comprises a nucleotide sequence encoding a doxycycline repressible promoter that has at least 90% identity to that of SEQ ID NO: 10. In some embodiments, the plasmid comprises a nucleotide sequence encoding a doxycycline repressible promoter that has at least 95% identity to that of SEQ ID NO: 10.

[0083] In some embodiments, the plasmid comprises a nucleotide sequence encoding an HIV LTR R5 region that has at least 85% identity to that of SEQ ID NO: 11. In some embodiments, the plasmid comprises a nucleotide sequence encoding an HIV LTR R5 region that has at least 90% identity to that of SEQ ID NO: 11. In some embodiments, the plasmid comprises a nucleotide sequence encoding an HIV LTR R5 region that has at least 95% identity to that of SEQ ID NO: 11.

[0084] In some embodiments, the plasmid comprises a nucleotide sequence encoding an HIV LTR U5 region that has at least 85% identity to that of SEQ ID NO: 12. In some embodiments, the plasmid comprises a nucleotide sequence encoding an HIV LTR U5 region that has at least 90% identity to that of SEQ ID NO: 12. In some embodiments, the plasmid comprises a nucleotide sequence encoding an HIV LTR U5 region that has at least 95% identity to that of SEQ ID NO: 12.

[0085] In some embodiments, the plasmid comprises a nucleotide sequence encoding a chromatin insulator that has at least 85% identity to that of SEQ ID NO: 13. In some embodiments, the plasmid comprises a nucleotide sequence encoding a chromatin insulator that has at least 90% identity to that of SEQ ID NO: 13.
embodiments, the plasmid comprises a nucleotide sequence encoding a chromatin insulator that has at least 95% identity to that of SEQ ID NO: 13.

[0086] In some embodiments, the plasmid comprises a nucleotide sequence encoding a beta-globin polyadenylation signal that has at least 85% identity to that of SEQ ID NO: 14. In some embodiments, the plasmid comprises a nucleotide sequence encoding a beta-globin polyadenylation signal that has at least 90% identity to that of SEQ ID NO: 14. In some embodiments, the plasmid comprises a nucleotide sequence encoding a beta-globin polyadenylation signal that has at least 95% identity to that of SEQ ID NO: 14.

[0087] In some embodiments, the plasmid comprises a nucleotide sequence that has at least 85% identity to that of SEQ ID NO: 15. In some embodiments, the plasmid comprises a nucleotide sequence that has at least 90% identity to that of SEQ ID NO: 15. In some embodiments, the plasmid comprises a nucleotide sequence that has at least 95% identity to that of SEQ ID NO: 15.

[0088] The disclosure provides lentiviral transfer vector plasmids incorporating a MCS for a variety of different restriction enzymes. According to certain embodiments of the disclosure, the MCS comprises a sequence having between about 20 and 40 nucleotides. In some embodiments, the MCS of the presently disclosed plasmid comprises at least two restriction enzyme cutting sites. In other embodiments, the MCS of the presently disclosed plasmid comprises at least three restriction enzyme cutting sites. In yet other embodiments, the MCS of the presently disclosed plasmid comprises between about 2 and about 10 restriction sites. In some embodiments, the restriction sites within the MCS are selected from the group consisting of BstBI, MluI, NotI, ClaI, ApaI, XhoI, XbaI, HpaI, NheI, Pael, NsiI, Sphi, Sma/Xma, Acel, BamHI, and Sphi, or any derivatives or analog thereof.

[0089] In some embodiments, the MCS region of the lentiviral transfer vector plasmid carries four unique restriction enzyme cutting sites which are believed to facilitate easy sub-cloning of a desired transgene cassette. In some embodiments, the multiple cloning site comprises the BstBI, MluI, NotI, and ClaI restriction endonuclease sites. In some embodiments, the nucleotide sequence encoding the multiple cloning site comprises the sequence of SEQ ID NO: 7, or a sequence having at
least 90% identity to that of SEQ ID NO: 7. There restriction site may be arranged in any order.

[0090] In some embodiments, the transfer plasmid comprises one or more additional restriction enzyme cutting sites flanking the vector backbone (see Fig. 11). Without wishing to be bound by any particular theory, it is believed that the additional flanking restriction enzyme cutting sites allow for the generation of a directional (a "head-to-tail") concatemeric array. In some embodiments, the restriction enzyme cutting sites are selected from SfiI and Bsu36I. In some embodiments, a lentiviral vector comprising one or more genes is derived from the plasmid.

[0091] In some embodiments, the lentiviral vector transfer plasmid comprises a nucleotide sequence having at least 80% identity to that of sequence of SEQ ID NO: 1. In other embodiments, the lentiviral vector transfer plasmid comprises a nucleotide sequence having at least 85% identity to that of sequence of SEQ ID NO: 1. In yet other embodiments, the lentiviral vector transfer plasmid comprises a nucleotide sequence having at least 90% identity to that of sequence of SEQ ID NO: 1. In further embodiments, the lentiviral vector transfer plasmid comprises a nucleotide sequence having at least 95% identity to that of sequence of SEQ ID NO: 1. In yet further embodiments, the lentiviral vector transfer plasmid comprises a nucleotide sequence having at least 97% identity to that of sequence of SEQ ID NO: 1. In some embodiments, the lentiviral vector transfer plasmid comprises the sequence of SEQ ID NO: 1. In some embodiments, the lentiviral vector transfer plasmid has a sequence that differs by not more than 100 nucleotides from the sequence set forth in SEQ ID NO: 1.

[0092] In some embodiments, the lentiviral vector transfer plasmid comprises a nucleotide sequence having at least 80% identity to that of sequence of SEQ ID NO: 2. In other embodiments, the lentiviral vector transfer plasmid comprises a nucleotide sequence having at least 85% identity to that of sequence of SEQ ID NO: 2. In yet other embodiments, the lentiviral vector transfer plasmid comprises a nucleotide sequence having at least 90% identity to that of sequence of SEQ ID NO: 2. In further embodiments, the lentiviral vector transfer plasmid comprises a nucleotide sequence having at least 95% identity to that of sequence of SEQ ID NO: 2. In yet further embodiments, the lentiviral vector transfer plasmid comprises a nucleotide sequence...
having at least 97% identity to that of sequence of SEQ ID NO: 2. In some embodiments, the lentiviral vector transfer plasmid comprises the sequence of SEQ ID NO: 2. In some embodiments, the lentiviral vector transfer plasmid has a sequence that differs by not more than 100 nucleotides from the sequence set forth in SEQ ID NO: 2.

[0093] In some embodiments, the lentiviral transfer vector plasmid is synthesized according to those methods known to those of skill in the art. For example, the plasmids may be synthesized using traditional restriction digestion and ligation techniques known to those of ordinary skill in the art. For example, a donor plasmid comprising the TL20c vector backbone may be subcloned into a pU57C recipient plasmid (e.g. such as those available commercially from Genescript), using standard digestion and ligation procedures known to those of ordinary skill in the art (see, for example, Sambrook et al. (1989) Molecular Cloning: A Laboratory Manual, 2nd Ed. Cold Spring Harbor, N.Y., the disclosure of which is hereby incorporated by reference herein in its entirety).

[0094] The present disclosure also includes a method of producing a lentiviral vector, e.g. LVsh5/C46. In one embodiment, the method comprises synthesizing a cDNA of a gene and cloning the synthesized cDNA into a restriction site of a plasmid, such as pUC57-TL20c. Genes may be inserted into an appropriate cloning site using techniques known to those of skill in the art. For example, a gene may be amplified by PCR and then cloned into a plasmid containing a desired promoter or gene-expression controlling element.

[0095] In some embodiments, and solely by way of example, the method comprises synthesizing a cDNA of a gene which expresses a protein capable of preventing HIV fusion into a cell or HIV replication; and then cloning the synthesized cDNA into a restriction site in a plasmid as disclosed herein.

[0096] **Generation of a Stable Producer Cell Line and Harvesting of Lentiviral Vectors Produced Therefrom**

[0097] In some embodiments of the present disclosure are methods of forming a stable producer cell line and harvesting lentiviral vectors produced from the generated stable producer cell line. With reference to FIG 2, the first step in producing a stable
producer cell line is to generate DNA fragments (10), such as from a lentiviral vector transfer plasmid and a second plasmid, such as an antibiotic resistance cassette plasmid. Following DNA fragment generation (10), the DNA is then used to form a concatemeric array (20). Subsequently, the concatemeric array is then introduced, such as by transfection, into a packaging cell line (30) (e.g. GPR, GPRG, GPRT, GPRG, GPRT-G or derivatives thereof packaging cell lines). Following introduction of the array (30) and subsequent transfection, clones are selected (40) and isolated (50) to generate the stable producer cell line (60). Vector supernatant comprising lentiviral vector may then be harvested.

**Concatemeric Array Formation and Purification**

A "concatemer" or "concatemeric array" (used interchangeably herein) (a long continuous DNA molecule that contains multiple copies of the same DNA sequences linked directly or indirectly in series) is generated and used in the transfection of the packaging cell line. In some embodiments, the concatemers are large arrays of linked vector genome expression cassettes, with antibiotic resistance cassettes interspersed therein.

With reference to FIG. 14, to form the concatemeric array, DNA fragments from a lentiviral transfer vector plasmid (step 100) and an antibiotic resistance cassette plasmid are generated (step 110). In some embodiments, the DNA fragments may be prepared by digesting each of the plasmids according to protocols known to those of ordinary skill in the art and then ligating the digested fragments. In some embodiments, electrophoresis and agarose gel are utilized to acquire the desired DNA fragments (step 120). In some embodiments, a DNA fragment concentration may be determined using a NanoDrop Spectrophotometer (step 130). A variety of strategies are available for ligating fragments of DNA, the choice of which depends on the nature of the termini of the DNA fragments and which choices can be readily made by the skilled artisan.

In some embodiments, the lentiviral transfer vector plasmid is based on pUC57-TL20c. In some embodiments, the antibiotic resistance cassette plasmid is driven by the PGK promoter. In some embodiments, the antibiotic resistance cassette plasmid comprises flanking sites for concatemerization with the lentivirus cassette in
the lentiviral transfer vector plasmid. In some embodiments, the antibiotic resistance cassette plasmid is PGK-ble (bleomycin resistance). In some embodiments, the PGK-ble plasmid comprises a nucleotide sequence having at least 90% identity to the sequence of SEQ ID NO: 9. In some embodiments, the concatemeric arrays is formed through the in vitro ligation of the DNA fragments from the lentiviral transfer vector plasmid and the PGK-ble plasmid.

[0102] FIG. 15 outlines the general steps used to form the concatemeric array. At step 200, generated DNA fragments are mixed and the volume of fragments in the ligation reaction are maximized to maintain the desired ratio (step 210). In some embodiments, a ratio of an amount of lentiviral transfer vector plasmid DNA to an amount of antibiotic resistance cassette plasmid DNA ranges from about 100:1 to about 1:100. In other embodiments, a ratio of an amount of lentiviral transfer vector plasmid DNA to an amount of antibiotic resistance cassette plasmid DNA ranges from about 50:1 to about 1:50. In yet other embodiments, a ratio of an amount of lentiviral transfer vector plasmid DNA to an amount of antibiotic resistance cassette plasmid DNA ranges from about 25:1 to about 1:25. In further embodiments, a ratio of an amount of lentiviral transfer vector plasmid DNA to an amount of antibiotic resistance cassette plasmid DNA ranges from about 10:1 to about 1:10.

[0103] In some embodiments, the concatemeric reaction mixture is incubated overnight at room temperature (step 220). Subsequently, the DNA fragment concentration for each sample may then be measured using a NanoDrop Spectrophotometer (step 230).

[0104] In some embodiments, a directional concatemeric array is formed and used in the transfection of a packing cell line. In some embodiments, the formation of the directional array is achieved by utilizing the restriction enzyme sites of the lentiviral transfer vector plasmid which flank the lentiviral vector backbone. In some embodiments, restriction digestion utilizes the restriction enzyme sites flanking the TL20c vector cassette and allows for the formation of nucleotide nonpalindromic overhangs, which can only be used to ligate from heat to tail. In some embodiments, directional ligation, according to the methods described herein, allow for the generation of a concatemeric array which comprises predominantly head-to-tail DNA products.
[0105] In some embodiments, the concatemeric array is formed according to the method set forth in Example 3 herein. Of course, the skilled artisan will recognize that the procedure provided in Example 3 may be adapted for the formation of a concatemeric array having different ratios of a first plasmid to a second plasmid and for transfer plasmids other than LVsh5/C46.

[0106] In some embodiments, the concatemeric array is purified by phenol-extraction and ethanol precipitation prior to transfection into a packing cell line. While this conventional technique is cheap and effective, however, the procedure is time consuming and may not yield reproducible yields. There is believed to be a risk of phenol/chloroform carry-over into the final sample. Moreover, the process is believed to involve hazardous chemicals and may generate toxic waste that must be disposed of with care and in accordance with hazardous waste guidelines.

[0107] Alternatively, in other embodiments, a silica-based method is used to purify the newly synthesized concatemeric array after ligation. This method is believed to provide a simple, reliable, fast, and convenient way for isolation of the high-quality transfection-grade concatemeric array. In some embodiments, the concatemeric array is purified using a DNeasy Mini spin column, available from Qiagen, such as using the procedure set forth in Example 6.

[0108] Transfection / Single Clone Isolation

[0109] Following purification of the concatemeric array, the array is then used to transfect packaging cell line cells. As used herein, the terms "transformation" and "transfection" are intended to refer to a variety of art-recognized techniques for introducing foreign nucleic acid (e.g., DNA or RNA) into cells. As will be evident from the examples provided herein, when a host cell permissive for production of lentiviral particles is transfected with the generated concatemeric array, the cell becomes a producer cell, i.e. a cell that produces infectious lentiviral particles.

[0110] In general, the concatemeric array or directional concatemeric array may be introduced into cells via conventional transfection techniques. With reference to FIG. 16, in some embodiments, cells are harvested and seeded 20-24 hours before transfection (step 300) and then transfected (step 320) with the concatemeric array
synthesized (step 310). A procedure for transfecting a packing cell line cell is provided as Example 4 herein.

[0111] One packaging cell line suitable for transfection with the formed concatemeric arrays is the GPR packaging cell line. The GPR line is an HIV-1-based packaging cell line derived from 293T/17 cells with the necessary viral components gagpol and rev (see, Throm et al., Efficient construction of producer cell lines for a SIN lentiviral vector for SCID-X1 gene therapy by concatemeric array transfection. Blood 113: 5104–5110, the disclosure of which is hereby incorporated by reference herein in its entirety).

[0112] Another packaging cell line suitable for transfection with the formed concatemeric arrays is a GPRG packaging cell line. In some embodiments, the GPRG packing cell line comprises gagpol, rev, and VSV-G.

[0113] Yet another packaging cell line suitable for transfection with the formed concatemeric arrays is a GPRT packaging cell line (gagpol, rev, and tat). GPRG and GPRT packaging cell lines and methods of forming the same are also disclosed by Throm et al., the disclosure of which is again hereby incorporated by reference herein in its entirety. Other suitable packaging cell lines (e.g. GPRT-G) are described by Wielgosz et al. "Generation of a lentiviral vector producer cell clone for human Wiskott-Aldrich syndrome gene therapy," Molecular Therapy—Methods & Clinical Development 2, Article number: 14063 (2015), the disclosure of which is hereby incorporated by reference herein in its entirety.

[0114] The skilled artisan will appreciate that other packaging cell lines suitable for use with the presently disclosed method may also be utilized. In some embodiments, other packaging cell lines may be derived from the GPR, GPRG, GPRT, or GPRT-G packaging cell lines. Without wishing to be bound by any particular theory, it is believed that the GPRT-G cell line has higher transduction efficiency in CD34+ cells (see Wielgosz). By "derived from," what is meant is a population of cells clonally descended from an individual cell and having some select qualities, such as the ability to produce active protein at a given titer, or the ability to proliferate to a particular density.
FIG. 17 illustrates the general process of selecting for transfected cells. In some embodiments, and after about 72 hours after transfection, GPRG cells are cultured with the selective media (zeocin and doxycycline) (step 400). The cells are then fed with selective medium (zeocin and doxycycline) every 3-4 days until the cell foci are identified (step 410). Subsequently, the cell lines are expanded and evaluated (step 420).

In some embodiments, following transfection a single foci selection/screening process is utilized to identify the single cell clones that have good manufacturing potential. According to this method, in some embodiments, selected cells are seeded sparsely in 150 × 25 mm dishes and allowed to expand and form discernible colonies for 2-3 weeks. The individual colonies can then be transferred to another smaller culture vessel for monoclonal expansion. This method is believed to be a cost-effective and frequently adopted technique; however, due to the nature limitations in the single foci selection technique, achieving a high probability of monoclonality of a good producing cell line may be challenging.

FIG. 18 illustrates single colony isolation. At step 500, flow cytometry is utilized to prepare the single cell sorting. The cells are then plated (step 510) in the conditioned culture media, and expanded (step 520).

In other embodiments, in order to generate a high titer lentiviral vector stable producer cell line, Fluorescence activated cell sorter (FACS) have been used to isolate single clones (see, e.g. FIG. 8). Conditioned medium, e.g. Zeocin (50ug/mL) and Doxycycline (1ng/mL), may also be added during a sorting process to increase cell attachment and viability, and promote colony formation. The use of conditioned growth media and the high throughput ability of FACS system is believed to enable the screening a large number of clones and thus is believed to increases the probability of finding high titer lentiviral vector producer clones.

In some embodiments, a clone with good growing rate and viral production ability is tested for stability over about 20 passages.

Induction of Producer Cell lines to Generate Virus
Following selection and expansion of the selected clones, the selected clones are induced to produce vector supernatants, and induction may be carried out according to the procedures known to those of ordinary skill in the art. In order to generate the lentiviral vectors produced by induced stable producer lines, the culture supernatant is harvested, in some embodiments, every day for up to 7 days. This production protocol can be easily utilized to produce a variety of test vectors at small scale. The repeated virus harvesting protocol can also increase the final yield of viral vectors. However, the daily harvest and media exchange is often not economical, and as an alternative to the of daily harvest, a new two-day harvest protocol has been devised. This new viral vector production protocol, described below, allows the same amount of viral vectors to be generated with less culture media consumptions.

FIG. 19 further illustrates the process of induction and evaluation. At step 600, the viral vectors are induced and then the spinoculation of 293T is conducted to determine transduction efficiency (step 610). The top three clones are screened (step 620) and expanded (step 630). The clones are then stored (e.g. under liquid nitrogen) (step 640).

Harvest every two days

Applicants have unexpectedly discovered that a two-day harvest allows for the generation of about the same amount of viral vectors as with a more traditional daily harvest, while also providing the benefit of requiring less culture medium.

In one embodiment is a first method of generating viral vectors from a two-day harvest according to the present invention, the first method comprising the following steps:

(1) Remove the old culture medium of the producer lines culture dish as completely as possible and wash the cells with 1×PBS.

(2) Add TrypLE™ Express Enzyme (1×) to the culture dish (available from ThermoFisher Scientific).

(3) Place in 37°C incubator for 2 minutes.
(4) Wash out the cells by adding D10 medium (no drug) and dissociate cell clusters into single cells by pipetting up and down (D10 media: Dulbecco's Modified Eagle Medium with high glucose, GlutaMAX™ Supplement and 10% (w/v) FBS and 1% (w/v) Pen/Strep).

(5) Centrifuge cells for 5 minutes at 4°C at 1200 rpm.

(6) Aspirate medium and gently suspend pellet in fresh D10 medium (no drug).

(7) Seed the cells about 95% confluent at culture dish (Roughly by plating 4 × 106 Cells/6-mm culture dish, Vectors Induction).

(8) The seeded cells were supplemented with fresh, pre-warm D10 medium after 24hr (Day 1 post-induction).

(9) Viral vectors can be first time harvested from the cells 48 hours after the first time media change (Day 3 post-induction).

(10) Add the fresh, pre-warm medium to the culture dish.

(11) Conduct the second time viral vector harvested 48 hours after the second time medium change (Day 5 post-induction).

(12) Add the fresh, pre-warm medium to the culture dish.

(13) Conduct the third time viral vector harvested 48 hours after the third time medium change (Day 7 post-induction).

Applicants have found that the viral titer can be yielded at second time viral vectors collection at Day 4 and Day 5 post-induction, as compared with more traditional methods where viral vectors could be collected seven days post induction.

In another embodiment is a second method of generating viral vectors from a two-day harvest according to the present invention, the first method comprising the following steps:

(1) Remove the media of the producer lines culture dish as completely as possible and wash the cells with 1×PBS.
(2) Gently Pipette 1×TrypLE Express onto the washed cell monolayer using 3 mL for 100 mm culture dish.
(3) Rotate flask to cover the monolayer with TrypLE Express.
(4) Return flask to the incubator and leave for 2 minutes.
(5) Gently tapped side of the flasks to release any remaining attached cells.
(6) Re-suspend the cells in 2 mL of the fresh D10 media (no antibiotics) and transfer to a 15 mL conical centrifuge Tube.
(7) Centrifuge cells for 5 minutes at 1200 rpm.
(8) Aspirate media and gently suspend pellet in 5 mL fresh D10 culture media (no antibiotics)
(9) Determine the cell counts by TC10™ Automated Cell Counter
(10) Seed the cells > 95% confluent at culture dish (by plating 4×10^6 Live Cells in 60-mm culture dish)
(11) The seeded cells were supplemented with fresh, warm-up D10 media daily (Every 24 hours).

Applicants have discovered that viral vectors could be harvested from the cells 48 hours post induction and that the highest viral titer could be yielded at 72 hours after induction. Applicants have again unexpectedly discovered that viral vectors could be harvested from days 2 - 4 post induction.

In some embodiments, the harvested vectors are purified through filtration. In some embodiments, the harvested vectors are characterized by determining viral titer, viral copy per cell genome, and p24 concentration.

Daily comparison of daily harvesting versus two-day harvesting is illustrated in Figure 5.

Example 1—Detailed Comparison of Self-Inactivating Lentiviral Vectors Produced by Transient Transfection and Vectors Produced by the Disclosed Stable Cell Line Method
The methods described herein were used to generate a stable cell line for the production of LVsh5/C46, a self-inactivating lentiviral vector (SIN-LV) encoding a short hairpin RNA (shRNA) for down-regulation of the HIV-1 co-receptor CCR5, in combination with the HIV-1 fusion inhibitor, C46. This LV, produced by transient transfection, is currently being evaluated in clinical trials in HIV-infected individuals. Here we conducted a comparative analysis of LVsh5/C46 produced by transient transfection and LVsh5/C46 produced using the methods described herein to support the application of this system for clinical manufacturing of LVsh5/C46 and other SIN-LVs.

Lentiviral vectors (LVs) were produced by calcium phosphate transfection in 293T cells using the 4-plasmid system (one transfer vector, two packaging vectors, and one envelope vector). Virus-containing media (VCM) was harvested 48h post-transfection and concentrated by ultracentrifugation through a 20% sucrose cushion.

For cell line production, producer cells were induced in media without doxycycline (Dox), and the VCM was harvested at 72h and similarly concentrated by ultracentrifugation. With reference to Table 1 and FIGs. 8 and 9, LVs produced by each method were compared based on particle titer and using three independent assays for gene transduction potency on 293T and the TF-1a T cell line. These included FACS assays for cell surface C46 expression and shRNA-mediated knockdown of CCR5 expression, as well as a qPCR assay for vector copy number (VCN) per host cell genome. For all assays, titer was determined over a range of vector dilutions to define a linear relationship. The qPCR assay utilized genomic DNA extracted from transduced cells, and detect the C46 transgene and a sequence from the endogenous β-globin gene. As such, C46 VCN could be normalized to cellular genome.
Table 1. Stable vs. Transient viral vector production.

<table>
<thead>
<tr>
<th>VCM²</th>
<th>Method</th>
<th>Titer 293T (TU/mL)</th>
<th>Titer TF-1a (TU/mL)</th>
<th>p24 (ng/mL)</th>
</tr>
</thead>
<tbody>
<tr>
<td>LVsh5/C46</td>
<td>Transient Transfection</td>
<td>$2.78 \times 10^8$</td>
<td>$2.64 \times 10^8$</td>
<td>5250</td>
</tr>
<tr>
<td>TL20sh5/C46</td>
<td>Stable Producer Cell</td>
<td>$1.40 \times 10^8$</td>
<td>$1.46 \times 10^8$</td>
<td>13430</td>
</tr>
</tbody>
</table>

1. Abbreviation: TU, Transduction Unit; VCM, virus-containing medium
2. VCM were concentrated 100-fold through a 20% sucrose cushion by ultracentrifugation

A higher concentration of p24 was observed in VCM produced by producer cell lines relative to transient transfection method. However, yield and potency of LVsh5/C46 produced using the two different systems was similar. Vectors were first evaluated for C46 titer by FACS using equal volumes of VCM. While vector produced by transient transfection had a modestly increased titer, when C46 titers were normalized and vector preparations were assed for gene transduction using the qPCR assay or via functional knock-down of CCR5, vector produced by the stable producer cell lines showed greater potency (see Table 2). Down-regulation of CCR5 expression and genomic C46 transgene (VCN) were each significantly higher in the target cells treated with LVsh5/C46 produced by the methods disclosed herein than treated with vector produced by transient transfection (see Table 3 and Figure 10).
Table 2. Analysis of C46 by qPCR in transduced cells

<table>
<thead>
<tr>
<th>Condition</th>
<th>MOI</th>
<th>C46 copies/cells</th>
</tr>
</thead>
<tbody>
<tr>
<td>no virus, negative ctrl.</td>
<td>-</td>
<td>ND</td>
</tr>
<tr>
<td>TF-1α², positive ctrl.</td>
<td>-</td>
<td>1.36 ± 0.58</td>
</tr>
<tr>
<td>Transient Transfection</td>
<td>1</td>
<td>5.34 ± 0.55</td>
</tr>
<tr>
<td>Stable Producer Line</td>
<td>1</td>
<td>10.7 ± 2.17</td>
</tr>
<tr>
<td>Transient Transfection</td>
<td>0.3</td>
<td>1.01 ± 0.30</td>
</tr>
<tr>
<td>Stable Producer Line</td>
<td>0.3</td>
<td>3.67 ± 0.66</td>
</tr>
</tbody>
</table>

1. Abbreviation: ND, Not Detected; MOI, Multiplicity of Infection
2. LVsh5/C46 single copy cell line
3. MOI based on C46 transduction titer
Table 3. Analysis of C46 by qPCR in transduced cells

<table>
<thead>
<tr>
<th>Condition</th>
<th>MOI$^3$</th>
<th>C46 copies/cells</th>
</tr>
</thead>
<tbody>
<tr>
<td>no virus, negative ctrl.</td>
<td>-</td>
<td>ND</td>
</tr>
<tr>
<td>TF-1a$^2$, positive ctrl.</td>
<td>-</td>
<td>1.36 ± 0.58</td>
</tr>
<tr>
<td>Transient Transfection</td>
<td>1</td>
<td>5.34 ± 0.55</td>
</tr>
<tr>
<td>Stable Producer Line</td>
<td>1</td>
<td>10.7 ± 2.17</td>
</tr>
<tr>
<td>Transient Transfection</td>
<td>0.3</td>
<td>1.01 ± 0.30</td>
</tr>
<tr>
<td>Stable Producer Line</td>
<td>0.3</td>
<td>3.67 ± 0.66</td>
</tr>
</tbody>
</table>

1. Abbreviation: ND, Not Detected; MOI, Multiplicity of Infection
2. LVsh5/C46 single copy cell line
3. MOI based on C46 transduction titer

[0160] Based on three independent assays, we demonstrate that the methods described herein provide a stable LV production system capable of generating similar quality and quantity of SIN-LVs compared to transient transfection methods. The higher CCR5 down-regulation efficacy and C46 VCN in transduced cells (normalized to C46 titer) indicate that LVsh5/C46 produced by producer cells has better potency than those vectors generated using the conventional 4-plasmid transient transfection. By removing the tedious transient transfection step, without wishing to be bound by any particular theory, it is believed that this production system can be easily adapted to cGMP conditions for the manufacture of clinical grade materials for use in humans.
Example 2–Development and Characterization of GPRG-Based Producer Cell Lines for the Bio-production of Lentiviral Vectors for HIV Gene Therapy

The GPRG cell line system has previously been established for the clinical production of self-inactivating lentiviral vectors (SIN-LVs). Here we sought to establish producer cell lines based on GPRG for the production of LVsh5/C46, a SIN-LV currently being assessed in the clinic for treatment of HIV-infected individuals. This vector encodes two viral entry inhibitors; sh5, a short hairpin RNA to the HIV coreceptor CCR5, and C46, a viral fusion inhibitor. We also sought to define the stability of GPRG packaging cell line, the GRPG-based LVsh5/C46 producer cell line, and LVsh5/C46 production following tetracycline induction as required for regulatory filling and clinical application of the GPRG system for bio-production of LVsh5/C46.

GPRG cells were cultured in D10 media with doxycycline (Dox) and puromycin (Puro). To generate LVsh5/C46 producer cells, GPRG cells were transfected with the transfer plasmid TL20-LVsh5/C46 and a Zeocin-resistance plasmid as a concatemeric array. Individual clones were evaluated for their ability to produce LVsh5/C46 vector and maintained in D10 media with Dox, Puro, and Zeocin. To assess the stability of the parental GPRG cell line for LV production, GPRG cells were transfected with transfer vector every 10 passages over a 3-month period (50+ total passages) (see FIGs. 3A and 3B). Virus-containing media (VCM) was harvested 48h post-transfection and vector titer was assessed by complementary gene transduction assays. To assess the stability of LV production from the stable producer cell clones, cells were induced in D10 media without Dox. VCM was harvested 72h after induction and titer was similarly assessed over a range of vector dilutions. To analyze the stability of VSV-G expression following induction after long-term passage, GPRG cells were induced by Dox withdraw and then stained using a biotin-conjugated anti-VSV-G antibody, followed by a secondary staining with Streptavidin-Phycoerythrin.

GPRG cells demonstrate stringent tetracycline-regulated expression of VSV-G. This packaging cell line was able to produce up to 107 LV transduction units (TU)/mL after transfection with the LV transfer vector and maintained high-level LV
production for more than 50 passages in continuous culture (see FIGs. 6A and 6B). By utilizing concatemeric array transfection, we demonstrate efficient construction of a producer cell line based on GPRG for the production of LVsh5C46. This cell line consistently generated titers above 106 TU/mL. Further increases in titer could be achieved by recloning and selection of secondary producer cell lines. Titers peaked 2 to 5 days post-induction. We also showed that the established stable producer cell lines could routinely maintain LVsh5/C46 production with titers exceeding 106 TU/mL during continuous culture exceeding 25 passages.

The GPRG cell line efficiently expressed VSV-G on cell surfaces upon the removal of Dox. It could also generate high titers LVs after transfection of transfer vector plasmids. Moreover, this cell line allowed the derivation of high-titer producer cell lines for SIN-LVs. Producer cell lines demonstrated stable vector production during prolonged culture, and evaluation of the adaptability to adapt vector production to serum-free and suspension culture systems has been explored (see FIGs. 7A and 7B).

Example 3—Protocol for the Generation of a Concatemeric Array

Step 1

Prepare 500mL of 1×TAE running buffer by combining 490mL of Deionized water with 10mL 50×TAE ((Tris-acetate-EDTA) buffer).

Make the 1% agarose gel by adding 1g of Agarose and 100 mL of 1×TAE buffer (Add 2 mL 50×TAE with 98 mL Autoclaved water) into a beaker and microwaving the mixture until there is no solid particles or bubbles (about 2.5 minutes).

Allow the mixture to cool for 3 minutes.

Add 10 μL of GelRed™ into the Agarose gel mixture and stir (available from Biotium).

Assemble the gel caster and gel comb. Pour the mixture into the gel mold and let it cool for 30 minutes (capacity: 60 μL for big comb)

Once the gel is cooled, fill the box with 1×TAE buffer until the gel is completely submerged.
[0174] Prepare the digest reaction mixture at room temperature to linearize the DNA.

[0175] Digest 25 μg of the vector plasmid with the restriction enzyme SfiI. In a separate reaction, digest the resistance cassette plasmid PGK-ble with PflMI (10 μg is more than enough).

<table>
<thead>
<tr>
<th>Component</th>
<th>The ble marker</th>
<th>Vector</th>
</tr>
</thead>
<tbody>
<tr>
<td>DNA Name</td>
<td>PGK-ble</td>
<td></td>
</tr>
<tr>
<td>10 FastDigest Green Buffer</td>
<td>5 μL</td>
<td>10 μL</td>
</tr>
<tr>
<td>Plasmid DNA</td>
<td>10 μg</td>
<td>25 μg</td>
</tr>
<tr>
<td>FastDigest Enzyme 1: PflMI</td>
<td>5 μL</td>
<td>0 μL</td>
</tr>
<tr>
<td>FastDigest Enzyme 2: SfiI</td>
<td>0 μL</td>
<td>5 μL</td>
</tr>
<tr>
<td>Water, nuclease-free</td>
<td>To 50</td>
<td>To 100</td>
</tr>
<tr>
<td>Total Volume</td>
<td>50 μL</td>
<td>100 μL</td>
</tr>
</tbody>
</table>

[0176] Mix gently and incubate at 37°C in a heat for 15 min.

[0177] Add 10 μL of GeneRuler 1kb plus DNA ladder mixture (2 μL DNA ladder + 8 μL nuclease-free water) and 50 μL of sample mix into the available slots.

[0178] Turn on the electrophoresis machine and run with the voltage of 150 V for 1 hour.

[0179] Transfer the gel into the UVP PhotoDoc-It Imaging system, and obtain the image of the result.

[0180] Download the gel pictures from the Eye-Fi website.
[0181] Determine the DNA concentration of each sample using the NanoDrop 2000 Spectrophotometer.

[0182] **Step 2**

[0183] Cut DNA bands out of the Agarose gel.

[0184] Add 3 volumes of Buffer QG to 1 volume of gel (Generally add 500 μL QG).

[0185] Incubate at 50°C for 10 minutes after the gel slice has dissolved completely.

[0186] Apply the sample to the QIAquick column, and centrifuge for 1 min at 17,900 rpm (available from Qiagen).

[0187] Discard flow-through and place QIAquick column back in the same collection tube.

[0188] Add 0.5 ml of Buffer QG to QIAquick column and centrifuge for 1 min.

[0189] Add 0.75 ml of Buffer PE to QIAquick column and centrifuge for 1 min.

[0190] Discard the flow-through and centrifuge the QIAquick column for an additional 1 min at 17,900 rpm.

[0191] Place QIAquick column into a clean 1.5 ml microcentrifuge tube.

[0192] To elute DNA, add 35 μL of Buffer EB to the center of the QIAquick membrane and centrifuge the column for 1 min at 17,900 rpm (Buffer EB is 10mM Tris-cl, pH 8.5).

[0193] Measure the DNA fragments concentration using the NanoDrop 2000 Spectrophotometer (Table 1; Using EB buffer for the Blank measurement)

[0194] **Step 3**

[0195] Set up the ligation reaction in a 1.7 mL Eppendorf microcentrifuge tube on ice.
Use the pre-constructed spreadsheet (Concatemeric Ligations.xlsx) to calculate the volumes of each fragment that needs to be mixed to create about a 25:1 molar ratio of vector to PGK-ble.

Maximize the volume of fragments in the ligation reaction and maintain the desired molar ratio.

The T4 DNA Ligase Buffer should be thawed and re-suspended at room temperature (T4 DNA Ligase Buffer comprises the following components: 50mM Tris-HCl, 10mM MgCl₂, 1mM ATP, 10mM DTT, pH 7.5).

Pipette the ligation reaction. In the above example, we used the 90 µL DNA mixture by adding 10 µL of 10X ligation buffer (NEB Quick Ligation kit), and 0.5 µL of Ligase enzyme (available from New England BioLabs).

Prepare the following reaction mixture containing at room temperature:

<table>
<thead>
<tr>
<th>Component</th>
<th>Vector</th>
</tr>
</thead>
<tbody>
<tr>
<td>The vector fragment</td>
<td></td>
</tr>
<tr>
<td>The ble-resistant fragment</td>
<td></td>
</tr>
<tr>
<td>10 × T4 DNA Ligase Buffer</td>
<td>10</td>
</tr>
<tr>
<td>T4 DNA Ligase</td>
<td>0.5</td>
</tr>
<tr>
<td>Water, nuclease-free</td>
<td>To 90</td>
</tr>
<tr>
<td>Total Volume</td>
<td>90</td>
</tr>
</tbody>
</table>

Mix gently by pipetting up and down.

Incubate at room temperature for overnight.

Step 4

The concatemeric array was harvested and purified prior to transfection into GPRG cells by the silica-based membrane (DNeasy Blood & Tissue Kit).
Pipet the concatemeric array mixture into the DNeasy Mini spin column placed in a 2 ml collection tube.

Centrifuge at 8000 × g for 1 min. Discard flow-through and collection tube.

Place the DNeasy Mini spin column in a new 2 ml collection tube (provided) (available from Qiagen).

Add 500 µL Buffer AW1, and centrifuge for 1 min at 8000 × g.

Discard flow-through and collection tube.

Place the DNeasy Mini spin column in a new 2 ml collection tube (provided)

Add 500 µL Buffer AW2, and centrifuge for 3 min at 20,000 × g to dry the DNeasy membrane.

Discard flow-through and collection tube.

Place the DNeasy Mini spin column in a clean 1.7 ml Eppendorf microcentrifuge tube

Add 200 µL Buffer AE directly onto the DNeasy membrane.

Incubate at room temperature for 4 min

Centrifuge for 1 min at 8000 × g to elute the DNA mixtures.

Repeat elution once

Measure the concatemeric DNA concentration by the NanoDrop Lite Spectrophotometer

Example 4 – Protocol for Generating Producer Cell Lines Using a Concatemeric Array

Passage the cells at least 4 times after thawing before using them in the viral vector production.

Ensure that cells are healthy and greater than 95% viable before vectors induction using Trypan Blue method (Trypan Blue is commonly used in dye exclusion
procedures for viable cell counting. This method is based on the principle that live (viable) cells do not take up certain dyes, whereas dead (non-viable) cells do. Staining facilitates the visualization of cell morphology.

[0222] Culture the desired quantity of GPRG cells

[0223] Subculture the cells at least two times daily passage before seed the cells.

[0224] The first day, remove the medium of the GPRG cell lines culture dish and wash the cells with 1×PBS.

[0225] Gently Pipette 1×TrypLE Express onto the washed cell monolayer using 3 ml for T75 flask or 1 mL for T25 Flask.

[0226] Rotate flask to cover the monolayer with TrypLE Express.

[0227] Return flask to the incubator and leave for 2 minutes.

[0228] Gently tapped side of the flasks to release any remaining attached cells.

[0229] Re-suspend the cells in 2 mL of the fresh D10 medium and transfer to a 15 mL conical centrifuge tube.

[0230] Centrifuge cells for 5 minutes at 1200 rpm.

[0231] Aspirate medium and gently suspend pellet in 5 mL fresh D10 culture medium with Doxycycline (1ng/mL)

[0232] Determine the cell counts by TC10™ Automated Cell Counter (Table 5)

[0233] Seed the cells 20-24 hours before the transfection at 80% confluent at culture dish (by plating 3.2×106 Live Cells in 60-mm culture dish with Doxycycline; Table 9)

[0234] Prepare the concatemeric arrays formation (see, for example, Example 3).

[0235] The second day, allow CalPhosTM Mammalian Transfection Kit to come to room temperature prior to the transfection (Table 7) (available From ClonTech).

[0236] Purify the concatemeric DNA and measure the concentration (the concatemeric array may be purified according to the methods described herein)
Prepare the transfection plasmid DNA table (Table 8.; 4 mL, 60 mm culture dish).

For each transfection, prepare Solution A and Solution B in the separate 15 mL conical centrifuge tube

Bubbling Solution B (2 × HBS) by Pipette-aids and add the Solution A (DNA mixture) drop by drop.

Incubate the transfection solution at room temperature for 15 min

Gently add the transfection solution to the culture dish.

Gently move plates back and forth to distribute transfection solution evenly.

Incubate plates at 37°C for 4 hours in a CO2 incubator.

Warm-up 5 mL fresh D10 media per 60 mm culture dish in a 37°C CO2 incubator

After 4 hours wash with 1 mL pre-warmed D10 and change with 4 mL pre-warmed fresh D10 medium

Incubate at the 5% CO2 37°C incubator

48 hours after concatemer transfection, harvest the transfected GPRG cells (Perform the Subculturing Cells).

Re-plate the cells in the T150 flask or the 30 mL, 150mm culture dish with fresh D10 media containing Zeocin (50ug/ml) and Doxycycline (1ng/mL)

Feed the cells with selective medium (Zeocin, 50ug/ml) with Doxycycline (1ng/mL) every 3-4 days until the cell foci are identified (usually observed within 1–2 weeks)

**Example 5—Description of cell lines and sequences used to generate the GPRG packaging cell line**

HEK-293T/17 are a sub-clone of HEK-293T. These cells stably express SV-40 T antigen, and a particular clone was selected specifically for its high
transfectability. A master cell bank based on HEK-293T/17 was generated (HEK-293T/17 MCB).

[0252] SFG-IC-HIVgp-Ppac2 is a gamma retroviral vector that expresses codon-optimized HIV gagpol under control of the CMV promoter, with puromycin resistance. The plasmid (pSFG-IC-HIVgp-Ppac2) that was used to make this vector was constructed using the following components:


[0254] (2) CMV enhancer/promoter driven codon optimized HIV NL4-3 gagpol gene;


[0256] Infection of the HEK-293T/17 MCB with the SFG-IC-HIVgp-Ppac2 retroviral vector produced the GP cell line.

[0257] SFG-tc-revco is a gamma retroviral vector that expresses codon-optimized HIV rev under control of the tetracycline responsive promoter. The plasmid used to produce this vector (pSFG-tc-revco) was constructed using the following components:

[0258] (1) The HIV rev gene based on the NL4-3 strain sequence as above, and

[0259] (2) pSFG tcLuc ECT3 (described above)

[0260] SFG-tTA is a gamma retroviral vector that expresses the chimeric transcriptional transactivator under control of the retroviral LTR (Lindemann, D., Patriquin, E., Feng, S., and Mulligan, R.C. Versatile retrovirus vector systems for regulated gene expression in vitro and in vivo. Mol. Med. 3, 466–476 (1997)). It is

[0261] Infection of the GP cell line with SFG-tc-revco and SFG-tTA produced the GPR cell line


[0263] Infection of the GPR cell line with SFG-tc-VSVG produced the GPRG cell line.


[0265] Example 6—CONCATEMERIC ARRAY PURIFICATION

[0266] The concatamer was harvested and purified prior to transfection into GPRG cells by the silica-based membrane (DNeasy Blood & Tissue Kit).

[0267] Pipet the concatemeric array mixture into the DNeasy Mini spin column placed in a 2 mL collection tube.

[0268] Centrifuge at 6000 × g for 1 min. Discard flow-through and collection tube

[0269] Place the DNeasy Mini spin column in a new 2 mL collection tube (provided)

[0270] Add 500 μLBuffer AW1, and centrifuge for 1 min at 6000 × g.

[0271] Discard flow-through and collection tube
[0272] Place the DNeasy Mini spin column in a new 2 ml collection tube (provided)

[0273] Add 500 µL Buffer AW2, and centrifuge for 3 min at 20,000 × g to dry the DNeasy membrane.


[0275] Place the DNeasy Mini spin column in a clean 1.7 mL Eppendorf microcentrifuge tube

[0276] Add 200 µL Buffer AE directly onto the DNeasy membrane.

[0277] Incubate at room temperature for 4 min

[0278] Centrifuge for 1 min at 6000 × g to elute the DNA mixtures.

[0279] Repeat elution once (add new elution buffer)

[0280] Measure the concatemeric DNA concentration by the NanoDrop Lite Spectrophotometer.

[0281] **Example 7- TL20-UbeGFP & Cal1-WPRE Producer Cell Line**

[0282] The Table which follows summarizes two producer cell lines that were synthesized according to the methods describes herein. Data relating to the TL20-Cal1-wpre and TL20-Unc-GFP vectors is illustrated further in FIGs. 20A, 20B, and 20C.
### Selection and Screening

<table>
<thead>
<tr>
<th></th>
<th>TL20c-Ubc-GFP</th>
<th>TL20c-Cal1-WPRE</th>
</tr>
</thead>
<tbody>
<tr>
<td>Method</td>
<td>Single Cell Sorting&lt;sup&gt;2&lt;/sup&gt;</td>
<td>Single Cell Sorting&lt;sup&gt;2&lt;/sup&gt;</td>
</tr>
<tr>
<td>Seed Cell Density</td>
<td>1 cell/well</td>
<td>1 cell/well</td>
</tr>
<tr>
<td>Culture Medium</td>
<td>Conditioned Medium</td>
<td>Conditioned Medium</td>
</tr>
<tr>
<td>Efficiency of clone formation</td>
<td>28/96</td>
<td>22/96</td>
</tr>
<tr>
<td>Complete Expanded</td>
<td>16</td>
<td>17</td>
</tr>
<tr>
<td>Evaluated clones</td>
<td>16</td>
<td>5</td>
</tr>
<tr>
<td>Polyclonal Vector Production</td>
<td>$5.77 \times 10^5$ TU/mL</td>
<td>$4.50 \times 10^5$ TU/mL</td>
</tr>
<tr>
<td>The productivity of the best clones</td>
<td>$1.36 \times 10^7$ TU/mL</td>
<td>$3.0 \times 10^6$ TU/mL</td>
</tr>
</tbody>
</table>

1. Abbreviation: TU, Transduction Unit
2. Performing single cell sorting by using flow cytometer at USC Flow Cytometry Core Facility
3. Conditioned Media: DMEM w GlutaMax; FBS (10% w/v); Pen/Strep (1% w/v); Doxycycline (1 ng/mL)

All publications mentioned in this specification are herein incorporated by reference in their entirety. It will be appreciated by persons skilled in the art that numerous variations and/or modifications may be made to the disclosure as shown in the specific embodiments without departing from the spirit or scope of the disclosure as broadly described. The present embodiments are, therefore, to be considered in all respects as illustrative and not restrictive.

Although the disclosure herein has been described with reference to particular embodiments, it is to be understood that these embodiments are merely illustrative of the principles and applications of the present disclosure. It is therefore to be understood that numerous modifications may be made to the illustrative embodiments and that other arrangements may be devised without departing from the spirit and scope of the present disclosure as defined by the appended claims.
CLAIMS

1. A recombinant plasmid comprising a nucleotide sequence having at least 90% identity to that of SEQ ID NO: 1.

2. The recombinant plasmid of claim 1, wherein the nucleotide sequence has at least 95% identity to that of sequence of SEQ ID NO: 1.

3. A recombinant plasmid comprising the nucleotide sequence of SEQ ID NO: 1.

4. A recombinant plasmid comprising a nucleotide sequence having at least 90% identity to that of SEQ NO: 2.

5. A recombinant plasmid having a sequence that differs by not more than 100 nucleotides from the sequence set forth in SEQ ID NO: 1.

6. The recombinant plasmid of claim 5, wherein the sequence differs by not more than 50 nucleotides from the sequence set forth in SEQ ID NO: 1.

7. A recombinant plasmid comprising between about 6500 nucleotides and about 6750 nucleotides, and wherein the plasmid comprises a sequence having at least 90% identity to that of SEQ ID NO: 2.

8. The recombinant plasmid of claim 7, wherein the plasmid comprises between about 6600 nucleotides and about 6700 nucleotides.

9. The recombinant plasmid of claim 7, wherein the plasmid comprises about 6611 nucleotides.

10. A recombinant plasmid as set forth in FIG. 11 as pUC57-TL20c.

11. A recombinant plasmid comprising a multiple cloning site consisting essentially of BstBI, MluI, NotI, and Clal restriction endonuclease sites.

12. The recombinant plasmid of claim 11, wherein the plasmid further comprises a nucleotide sequence encoding a packaging signal; a nucleotide sequence encoding a central polypurine tract; a nucleotide sequence encoding a Rev response element; and a nucleotide sequence encoding a self-inactivating long terminal repeat.

13. A recombinant plasmid comprising (a) a nucleotide sequence encoding a packaging signal; (b) a nucleotide sequence encoding a central polypurine tract (cPPT); (c) a nucleotide sequence encoding a Rev response element; (d) a nucleotide sequence encoding a self-inactivating long terminal repeat; and (e) a
nucleotide sequence encoding a multiple cloning site having restriction sites for
the enzymes BstBI, Mlu I, Not I, and Cla I.
14. The recombinant plasmid of claim 13, where the nucleotide sequence encoding
the packing signal comprises the sequence of SEQ ID NO: 3.
15. The recombinant plasmid of claim 13, wherein the nucleotide sequence
encoding the central polypurine tract (cPPT) comprises the sequence of SEQ ID
NO: 4.
16. The recombinant plasmid of claim 13, wherein the nucleotide sequence
encoding the Rev response element comprises the sequence of SEQ ID NO: 5.
17. The recombinant plasmid of claim 13, wherein the nucleotide sequence
encoding the self-inactivating long terminal repeat comprises the sequence of
SEQ ID NO: 6.
18. The recombinant plasmid of claim 13, wherein the nucleotide sequence
encoding the multiple cloning site comprises the sequence of SEQ ID NO: 7.
19. A recombinant plasmid comprising (a) a packaging sequence, the packaging
sequence present from about nucleotide 762 to about nucleotide 1104 of a
nucleotide sequence of the plasmid; (b) a central polypurine tract, the central
polypurine tract present from about nucleotide 1121 to about nucleotide 1597 of
the plasmid nucleotide sequence; (c) a Rev response element, the Rev response
element present from about nucleotide 1598 to about nucleotide 2366 of the
plasmid nucleotide sequence; (d) a self-inactivating long terminal repeat, the
self-inactivating long terminal repeat present from about nucleotide 409 to
about nucleotide 589 of the plasmid nucleotide sequence; and (e) a multiple
cloning site, the multiple cloning site present from about nucleotide 2376 to
about nucleotide 2400 of the plasmid nucleotide sequence.
20. The recombinant plasmid of claim 19, wherein the plasmid nucleotide sequence
comprises a sequence having at least 90% identity to that of SEQ ID NO: 1.
21. The recombinant plasmid of claim 19, where a nucleotide sequence encoding
the packing signal comprises the sequence of SEQ ID NO: 3.
22. The recombinant plasmid of claim 19, wherein a nucleotide sequence encoding
the central polypurine tract (cPPT) comprises the sequence of SEQ ID NO: 4.
23. The recombinant plasmid of claim 19, wherein a nucleotide sequence encoding the Rev response element comprises the sequence of SEQ ID NO: 5.

24. The recombinant plasmid of claim 19, wherein a nucleotide sequence encoding the self-inactivating long terminal repeat comprises the sequence of SEQ ID NO: 6.

25. The recombinant plasmid of claim 19, wherein a nucleotide sequence encoding the multiple cloning site comprises the sequence of SEQ ID NO: 7.

26. A recombinant plasmid comprising a multiple cloning site comprising BstBI, MluI, NotI, and Clal restriction endonuclease sites, and wherein the plasmid comprises a nucleotide sequence having at least 80% identity to that of SEQ ID NO: 1.

27. A recombinant plasmid comprising a nucleotide sequence encoding a vector cassette having at least 95% identity to that of SEQ ID NO: 2, and wherein the vector cassette is flanked by at least two restriction endonuclease sites, the at least two restriction endonuclease sites independently selected from the group consisting of sfI and Bsu36I.

28. A recombinant plasmid comprising a nucleotide sequence encoding a vector cassette having at least 90% identity to that of SEQ ID NO: 2, the vector cassette comprising a multiple cloning site having BstBI, MluI, NotI, and Clal restriction endonuclease sites, wherein the plasmid further comprises a tetracycline repressible promoter upstream of the vector cassette.

29. A cell comprising the plasmid of any of claims 1 to 28.

30. A kit comprising the plasmid of any of claims 1 to 28 and a bleomycin resistance (ble) cassette.

31. A method of producing a stable producer cell line comprising: (a) synthesizing a lentiviral vector by cloning one or more genes into the plasmid of any of claims 1 to 28; (b) generating DNA fragments from the synthesized lentiviral vector; (c) forming a concatemeric array from the generated DNA fragments from the synthesized lentiviral vector and from DNA fragments from an antibiotic resistance cassette plasmid; (d) transfecting a GPR, GPRG, GPR, GPRGT, GPR-T-G packing cell line or a derivative thereof with the formed concatemeric array; and (e) isolating the stable producer cell line.
32. A method of producing a stable producer cell line comprising: (a) synthesizing a lentiviral vector which encodes a short hairpin RNA for down-regulation of an HIV-1 co-receptor and which encodes an HIV-1 fusion inhibitor, the lentiviral vector synthesized by cloning cDNA encoding both the short hairpin RNA and the fusion inhibitor into the plasmid of any of claims 1 to 28; (b) generating DNA fragments from the synthesized lentiviral vector; (c) forming a concatemeric array from the generated DNA fragments from the synthesized lentiviral vector and from DNA fragments from an antibiotic resistance cassette plasmid; (d) transfecting a GPR, GPRG, GPRT, GPRGT, GPRT-G packing cell line or a derivative thereof with the formed concatemeric array; and (e) isolating the stable producer cell line.

33. A method of harvesting vector supernatant from a stable producer cell line, wherein the vector supernatant is harvested every about 40 hours to about 56 hours.

34. A method of harvesting vector supernatant comprising LVsh5/C46 lentiviral vector, wherein the vector supernatant is harvested every about 40 hours to about 56 hours.

35. A stable producer cell line suitable for producing LVsh5/C46.

36. The stable producer cell line of claim 35, wherein the stable producer cell line is based on the GPRG packaging cell line.

37. The stable producer cell line of claim 35, wherein the stable producer cell line is based on the GPRT packaging cell line.

38. The stable producer cell line of claim 35, wherein the stable producer cell line is based on the GPR packaging cell line.

39. A stable producer cell line suitable for producing a self-inactivating lentiviral vector, the lentiviral vector comprising a nucleotide sequence having at least 90% identity to that of SEQ ID NO: 8.

40. The stable producer cell line of claim 39, wherein the stable producer cell line is based on the GPRG packaging cell line.

41. The stable producer cell line of claim 39, wherein the stable producer cell line is based on the GPRT packaging cell line.
42. The stable producer cell line of claim 39, wherein the stable producer cell line is based on the GPR packaging cell line.

43. A concatemeric array comprising DNA fragments from a first plasmid and a second plasmid; the first plasmid derived from pUC57-TL20c; the second plasmid comprising a bleomycin antibiotic resistance cassette; wherein a ratio of the DNA fragments from the first plasmid to the second plasmid ranges from about 25:1 to about 1:25.

44. A stable producer cell line, the stable producer cell line generated by transfecting a packaging cell line selected from the group consisting of a GPR, GPRG, GPRT, GPRGT and GPRT-G packing cell line or a derivative thereof with the concatemeric array of claim 43.

45. The stable producer cell line of claim 44, wherein the stable producer cell line produces LVsh5/C46.

46. The stable producer cell line of claim 45, wherein the LVsh5/C46 is capable of being harvested every about 40 hours to about 56 hours.

47. An isolated vector having the plasmid map of FIG. 11 which comprises a multiple cloning site that comprises the nucleotide sequence set forth in SEQ ID NO: 7.

48. A method for making a plasmid vector comprising cleaving the plasmid of any of claims 1 to 28 and ligating the ends of the cleaved plasmid to compatible ends of an introduced polynucleotide.

49. The method of claim 48, wherein the introduced polynucleotide encodes at least one of a short hairpin RNA for down-regulation of an HIV-1 co-receptor or encodes an HIV-1 fusion inhibitor.

50. A kit comprising (a) the lentiviral transfer vector plasmid of any of claims 1 to 28; and (b) packaging cell line cells.

51. The kit of claim 50, wherein the packaging cell line cells are selected from the group consisting of GPR, GPRG, GPRT, GPRGT, and a derivative thereof.

52. The kit of claim 50, further comprising a bleomycin resistance (ble) cassette.

53. A lentiviral vector derived from the plasmid of any of claims 1-28.

54. The lentiviral vector of claim 53, wherein the lentiviral vector comprises at least one additional nucleotide sequence.
55. The lentiviral vector of claim 54, wherein the at least one additional nucleotide sequence is selected from the group consisting of a nucleotide sequence which encodes a short hairpin RNA for down-regulation of an HIV-1 co-receptor and a nucleotide sequence which encodes an HIV-1 fusion inhibitor.

56. The lentiviral vector of claim 55, wherein the lentiviral vector is LVsh5/C46.

57. The lentiviral vector of claim 55, wherein the lentiviral vector comprises a sequence having at least 95% identity to that of SEQ ID NO: 8.

58. A cell comprising the lentiviral vector of any of claims 53 to 57.

59. A pharmaceutical composition comprising the lentiviral vector of any of claims 53-57 and a carrier.
FIG. 1

- Transfection
- Producer cell line

Counts range from 1.00E+06 to 3.50E+06.
Generate DNA fragments

Synthesize a concatemer array

Introduce the concatemer array into GPRG cells

Select for transfected clones

Perform the single colony isolation

Evaluate viral production of each cell clone

FIG. 2
FIG. 4
FIG. 5
FIG. 6

A

VSV-G Density (MFU)

-Dox
+Dox
+Dox

no stained
SAVE-Cys only
SAVE-Cys only
SAVE-Cys only
SAVE-Cys only
SAVE-Cys only
SAVE-Cys only

alSVG-Bio + SAV Cys only
alSVG-Bio + SAV Cys only
alSVG-Bio + SAV Cys only
alSVG-Bio + SAV Cys only
alSVG-Bio + SAV Cys only
alSVG-Bio + SAV Cys only
alSVG-Bio + SAV Cys only

293T GPRG

B

Vector Titer (10^6 IU/mL)

100
10
1

8 11 21 32 42 51

Passage Times
FIG. 7
FIG. 9

qPCR C46 copies/cell

moi=1

moi=0.3

FIG. 11
FIG. 13

Vector titer (TU/mL)

Days after removal of dox

With serum
Serum-free
Digest the vector plasmid with the restriction enzyme SfiI

Digest the resistance cassette plasmid PGK-ble with the restriction enzyme PflMI

Utilize electrophoresis and agarose gel to acquire the desired DNA fragments

Determine the DNA fragment concentration of each sample using the NanoDrop Spectrophotometer

FIG. 14
Mix the generated DNA fragments to create a 25:1 molar ratio of vector to PGK-ble

Maximize the volume of fragments in the ligation reaction and maintain the desired molar ratio

Prepare the concatemeric reaction mixture incubate at room temperature for overnight

Determine the DNA fragments concentration of each sample using the NanoDrop Spectrophotometer

FIG. 15
16 / 21

Harvest and seed the cells 20-24 hours before the transfection

Prepare the concatemeric array formation

Transfect GPRG cell with the concatemeric DNA by CalPhos™ Mammalian Transfection Kit

FIG. 16
72 hours after transfection, culture the transfected GPRG cells with the selective media (zeocin and doxycycline)

Feed the cells with the selective medium (with zeocin and doxycycline) every 3-4 days until the cell foci are identified

Expand and polyclonal cell lines and evaluate vectors production ability

FIG. 17
Prepare the single cell sorting by the flow cytometry

Plate the sorted cells in the conditioned culture media with doxycycline

Expand the selected single colonies and store colonies at about 80°C

FIG. 18
Induce the viral vectors production of each clone

Conduct the spinoculation of 293T to determine the transduction efficiency

Screen the top three relatively fast growth and best titer clones

Expand the selected clones and generated R&D master and working cell banks

Storage at the liquid nitrogen tank

FIG. 19
FIG. 20A

FIG. 20B
FIG. 20C
Generate DNA fragments

Synthesize a concatemer array

Introduce the concatemer array into GPRG cells

Select for transfected clones

Perform the single colony isolation

Evaluate viral production of each cell clone

FIG. 2