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DESCRIPTION

TECHNICAL FIELD

[0001] The present disclosure is in the field of genome engineering, particularly targeted modification of the genome of a hematopoietic cell.

BACKGROUND

[0002] When one considers that genome sequencing efforts have revealed that the human genome contains between 20,000 and 25,000 genes, but fewer than 2000 transcriptional regulators, it becomes clear that a number of factors must interact to control gene expression in all its various temporal, developmental and tissue specific manifestations. Expression of genes is controlled by a highly complex mixture of general and specific transcriptional regulators that interact with DNA elements. These DNA elements comprise both local DNA elements such as the core promoter and its associated transcription factor binding sites as well as distal elements such as enhancers, silencers, insulators and locus control regions (LCRs) (see Matson et al. (2006) Ann Rev Genome Hum Genet 7: 29-50).

[0003] Enhancer elements were first identified in the SV40 viral genome, and then found in the human immunoglobulin heavy chain locus. Now known to play regulatory roles in the expression of many genes, enhancers appear to mainly influence temporal and spatial patterns of gene expression. It has also been found that enhancers can function to regulate expression at large distances from the core promoter of the targeted gene, and are not dependent on any specific sequence orientation with respect to the promoter. Enhancers can be located several hundred kilobases upstream or downstream of a core promoter region, where they can be located in an intron sequence, or even beyond the 3' end of a gene.

[0004] Various methods and compositions for targeted cleavage of genomic DNA have been described. Such targeted cleavage events can be used, for example, to induce targeted mutagenesis, induce targeted deletions of cellular DNA sequences, and facilitate targeted recombination at a predetermined chromosomal locus. See, e.g., U.S. Patent Nos. 9,255,250; 9,200,266; 9,045,763; 9,005,973; 9,150,847; 8,956,828; 8,945,868; 8,703,489; 8,586,526; 6,534,261; 6,599,692; 6,503,717; 6,689,558; 7,067,317; 7,262,054; 7,888,121; 7,972,854; 7,914,796; 7,951,925; 8,110,379; 8,409,861; U.S. Patent Publication Nos. 2003/0232410; 2005/0208489; 2005/0026157; 2005/0064474; 2006/0063231; 2008/0159996; 2010/00218264; 2012/0017290; 2011/0265198; 2013/0137104; 2013/0122591; 2013/0177983; 2013/0196373; 2015/0056705 and 2015/0335708.

[0005] These methods often involve the use of engineered cleavage systems to induce a double strand break (DSB) or a nick in a target DNA sequence such that repair of the break by an error born process such as non-homologous end joining (NHEJ), non-homology directed end capture of donors or repair using a repair template (homology directed repair or HDR) can result in the knock out of a gene or the insertion of a sequence of interest (targeted integration). See, e.g., U.S. Patent Nos. 9,045,763; 9,200,266; 9,005,973; and 8,703,489. These techniques can also be used to introduce site specific changes in the genome sequence through use of a donor oligonucleotide, including the introduction of specific deletions of genomic regions, or of specific point mutations or localized alterations (also known as gene correction). Cleavage can occur through the use of specific nucleases such as engineered zinc finger nucleases (ZFN), transcription-activator like effector nucleases (TALENs), or using the CRISPR/Cas system with an engineered crRNA/tracr RNA ('single guide RNA') to guide specific cleavage. Further, targeted nucleases are being developed based on the Argonaute system (e.g., from *T. thermophilus*, known as 'TtAgo', see Swarts et al. (2014) Nature 507(7491):

258-261), which also may have the potential for uses in genome editing and gene therapy.

[0006] Red blood cells (RBCs), or erythrocytes, are the major cellular component of blood and account for one quarter of the cells in a human. Mature RBCs lack a nucleus and many other organelles and are full of hemoglobin, a metalloprotein that functions to carry oxygen from the lungs to the tissues as well as carry carbon dioxide out of the tissues and back to the lungs for removal. This protein makes up approximately 97% of the dry weight of RBCs and it increases the oxygen carrying ability of blood by about seventy-fold. Hemoglobin is a heterotetramer comprising two alpha (α)-like globin chains and two beta (β)-like globin chains and 4 heme groups. In adults, the $\alpha_2\beta_2$ tetramer is referred to as Hemoglobin A (HbA) or adult hemoglobin. Typically, the alpha and beta globin chains are synthesized in an approximate 1:1 ratio and this ratio seems to be critical in terms of hemoglobin and RBC stabilization. In a developing fetus, a different form of hemoglobin, fetal hemoglobin (HbF), is produced which has a higher binding affinity for oxygen than Hemoglobin A such that oxygen can be delivered to the baby's system via the mother's blood stream. There are two genes that encode fetal globin that are very similar in sequence and are termed HBG1 (also referred to as Ggamma) and HBG2 (Agamma), based on their order of arrangement in the beta globin gene locus. Like adult hemoglobin, fetal hemoglobin protein contains two α globin chains, but in place of the adult β -globin chains, it has two fetal gamma (γ)-globin chains (*i.e.*, fetal hemoglobin is $\alpha_2\gamma_2$). At approximately 30 weeks of gestation, the synthesis of gamma globin in the fetus starts to drop, while the production of beta globin increases. By approximately 10 months of age, the newborn's hemoglobin is nearly all $\alpha_2\beta_2$ although some HbF persists into adulthood (approximately 1-3% of total hemoglobin). The regulation of the switch from production of gamma- to beta-globin is quite complex, and primarily involves a down-regulation of gamma globin transcription with a simultaneous up-regulation of beta globin transcription.

[0007] Genetic defects in the sequences encoding the hemoglobin chains can be responsible for a group of diseases known as hemoglobinopathies that include sickle cell anemia and the alpha and beta thalassemias. In the majority of patients with hemoglobinopathies, the genes encoding gamma globin remain present, but expression is relatively low due to normal gene repression occurring around parturition as described above.

[0008] It is estimated that 1 in 5000 people in the U.S. have sickle cell disease (SCD), mostly in people of sub-Saharan Africa descent (Roseff (2009) Immunohematology 25(2):67). There appears to be a benefit for heterozygous carriers of the sickle cell mutation due to protection against malaria, so this trait may have been positively selected over time, such that it is estimated that in sub-Saharan Africa, up to 28% of the population has the sickle cell trait (Elguero et al. (2015) PNAS USA 112 (22): 7051). Sickle cell disease is caused by a mutation in the β globin gene as a consequence of a valine substitution for glutamic acid at amino acid #6 (a GAG to GTG mutation at the DNA level), where the resultant hemoglobin is referred to as "hemoglobin S" or "HbS." Under lower oxygen conditions, a conformational shift in the deoxy form of HbS exposes a hydrophobic patch on the protein between the E and F helices. The hydrophobic residues of the valine at position 6 of the beta chain in hemoglobin are able to associate with the hydrophobic patch, causing HbS molecules to aggregate and form fibrous precipitates. These aggregates in turn cause the abnormality or 'sickling' of the RBCs, resulting in a loss of cell flexibility. The sickling RBCs are no longer able to squeeze into the capillary beds and can result in vaso-occlusive crisis in sickle cell patients. In addition, sickled RBCs are more fragile than normal RBCs, and tend towards hemolysis, eventually leading to anemia in the patient.

[0009] Treatment and management of sickle cell patients is a life-long proposition involving antibiotic treatment, pain management and transfusions during acute episodes. One approach is the use of hydroxyurea, which exerts its effects in part by increasing the production of gamma globin. Long term side effects of chronic hydroxyurea therapy are still unknown, however, and treatment gives unwanted side effects that lead to low patient compliance, and has variable efficacy from patient to patient (Brandow and Panepinto (2011) Am J Hematol 86(9):804-806). Despite an increase in the efficacy of sickle cell treatments, the life expectancy of patients is still only in the mid to late 50's and the associated morbidities of the disease have a profound impact on a patient's quality of life.

[0010] Thalassemias are also diseases relating to hemoglobin and typically involve a reduced expression of globin chains. This can occur through mutations in the regulatory regions of the genes or from a mutation in a globin coding sequence that results in reduced expression or reduced levels or functional globin protein. Alpha thalassemias, caused by mutations in the alpha globin locus, are mainly associated with people of Western Africa and South Asian descent, and may confer malarial resistance. Beta thalassemia, caused by mutations in the beta globin locus, is mainly associated with people of Mediterranean descent, typically from Greece and the coastal areas of Turkey and Italy. In thalassemia minor, only one of the β globin alleles bears a mutation. Individuals will suffer from microcytic anemia, and detection usually involves lower than normal mean corpuscular volume ($<80\text{fL}$). The alleles of subjects with thalassemia minor are β^+/β or β^0/β (where ' β^+ ' refers to alleles that allow some amount of β chain formation to occur, ' β ' refers to wild type β globin alleles, and ' β^0 ' refers to β globin mutations associated with a complete absence of beta-globin expression). Thalassemia intermedia subjects can often manage a normal life but may need occasional transfusions, especially at times of illness or pregnancy, depending on the severity of their anemia. These patient's alleles can be β^+/β^+ or β^0/β^+ . Thalassemia major occurs when both alleles have thalassemia mutations. This is severely microcytic and hypochromic anemia. Untreated, it causes anemia, splenomegaly and severe bone deformities and progresses to death before age 20. Treatment consists of periodic blood transfusion; splenectomy for splenomegaly and chelation of transfusion-caused iron overload. Bone marrow transplants are also being used for treatment of people with severe thalassemias if an appropriate donor can be identified, but this procedure can have significant risks.

[0011] One approach that has been proposed for the treatment of both SCD and beta thalassemias is to increase the expression of gamma globin with the aim to have HbF functionally replace the aberrant adult hemoglobin. As mentioned above, treatment of SCD patients with hydroxyurea is thought to be successful in part due to its effect on increasing gamma globin expression. The first group of compounds discovered to affect gamma globin expression were cytotoxic drugs. The ability to cause *de novo* synthesis of gamma-globin by pharmacological manipulation was first shown using 5-azacytidine in experimental animals (DeSimone (1982) Proc Nat'l Acad Sci USA 79(14):4428-31). Subsequent studies confirmed the ability of 5-azacytidine to increase HbF in patients with β -thalassemia and sickle cell disease (Ley, et al., (1982) N. Engl. J. Medicine, 307: 1469-1475, and Ley, et al., (1983) Blood 62: 370-380). In addition, short chain fatty acids (e.g. butyrate and derivatives) have been shown in experimental systems to increase HbF (Constantoulakis et al., (1988) Blood 72(6):1961-1967). Also, there is a segment of the human population with a condition known as 'Hereditary Persistence of Fetal Hemoglobin' (HPFH) where elevated amounts of HbF persist in adulthood (10-40% in HPFH heterozygotes (see Thein et al. (2009) Hum. Mol. Genet 18 (R2): R216-R223). This is a rare condition, but in the absence of any associated beta globin abnormalities, is not associated with any significant clinical manifestations, even when 100% of the individual's hemoglobin is HbF. When individuals that have a beta thalassemia also have co-incident HPFH, the expression of HbF can lessen the severity of the disease (Potoka and Gladwin (2015) Am J Physiol Lung Cell Mol Physiol. 308(4): L314-L324). Further, the severity of the natural course of sickle cell disease can vary significantly from patient to patient, and this variability, in part, can be traced to the fact that some individuals with milder disease express higher levels of HbF.

[0012] One approach to increase the expression of HbF involves identification of genes whose products play a role in the regulation of gamma globin expression. One such gene is BCL11A, first identified because of its role in lymphocyte development. BCL11A encodes a zinc finger protein that is thought to be involved in the developmental stage-specific regulation of gamma globin expression. BCL11A is expressed in adult erythroid precursor cells and down-regulation of its expression leads to an increase in gamma globin expression. In addition, the splicing of the BCL11A mRNA is developmentally regulated. In embryonic cells, the shorter BCL11A mRNA variants, known as BCL11A-S and BCL11A-XS are primarily expressed, while in adult cells, the longer BCL11A-L and BCL11A-XL mRNA variants are predominantly expressed. See, Sankaran et al. (2008) Science 322 p. 1839. The BCL11A protein appears to interact with the beta globin locus to alter its conformation and thus its expression at different developmental stages. Use of an inhibitory RNA targeted to

the BCL11A gene has been proposed (see, e.g., U.S. Patent Publication No. 2011/0182867) but this technology has several potential drawbacks, namely that complete knock down may not be achieved, delivery of such RNAs may be problematic and the RNAs must be present continuously, requiring multiple treatments for life.

[0013] Targeting of BCL11A enhancer sequences provides a mechanism for increasing HbF. See, e.g., U.S. Patent Publication No. 2015/0132269 and PCT Publication No. WO 2016/183298. Genome-wide association studies have identified a set of genetic variations at the BCL11A gene locus that are associated with increased HbF levels. These variations are a collection of small nucleotide polymorphisms (SNP) found in non-coding regions of BCL11A that function as a stage-specific, lineage-restricted enhancer region. Further investigation revealed that this BCL11A enhancer is required in erythroid cells for BCL11A expression, but is not required for its expression in B cells (see Bauer et al. (2013) Science 343:253-257). The enhancer region was found within intron 2 of the BCL11A gene, and three areas of DNaseI hypersensitivity (often indicative of a chromatin state that is associated with regulatory potential) in intron 2 were identified. These three areas were identified as "+62", "+58" and "+55" in accordance with the distance in kilobases from the transcription start site of BCL11A. These enhancer regions are roughly 350 (+55); 550 (+58); and 350 (+62) nucleotides in length (Bauer 2013, *ibid*).

[0014] When developing a nuclease for use in therapeutic treatments of humans, it is essential that the nuclease have the utmost safety characteristics. Specifically, the nucleases must have very low levels of off-target cleavage. Significant numbers of double strand cuts in locations other than the user-specified target can lead to repression of off-target genes, and in rare instances, the occurrence of chromosomal translocations (see Hoban and Bauer (2016) Blood, 127(21):2525-2535 and Tsang et al. (2017) Nature Methods, in press). Improvements in specificity can be achieved by eliminating non-specific interactions between the engineered nuclease and the genomic DNA (see U.S. Provisional Patent Application Nos. 62/378,978 and 62/443,981).

[0015] Thus, there remains a need for additional highly specific methods and compositions for the alteration of BCL11A gene expression, for example to treat hemoglobinopathies such as sickle cell disease and beta thalassemia.

SUMMARY

[0016] Described herein are highly specific compositions and methods for use in gene therapy and genome engineering. Specifically, the methods and compositions described relate to inactivating (e.g., by completely or partially abolishing its expression) a BCL11A gene, for example, a gene that acts as regulator of one or more additional genes. In particular, described are methods and compositions for interfering with enhancer function in a BCL11A gene to diminish or knock out its activity in specific cell lineages (e.g., erythroid). Additionally, described are methods and compositions for interfering with BCL11A enhancer functions wherein the enhancer sequences are not located within the coding sequences of the BCL11A gene, and wherein the reagents provided exhibit highly specific activity. The resulting down-regulation of the BCL11A gene in these circumstances results in increased expression of gamma globin, and the number of off-target cleavage events is reduced. The invention provides a zinc finger nuclease (ZFN) comprising the amino acid sequence as shown in SEQ ID NO:29 and the amino acid sequence as shown in SEQ ID NO:31.

The invention also provides one or more polynucleotides encoding one or more zinc finger nucleases according to the invention, optionally wherein the one or more polynucleotides is(are) mRNA.

The invention further provides one or more vectors comprising sequences encoding one or more zinc finger nucleases of the invention.

The invention provides a cell comprising the zinc finger nuclease of the invention, the one or more polynucleotides of the invention or the one or more vectors of the invention.

The invention also provides a cell or cell line produced from the cell of the invention, comprising the zinc finger

nuclease of the invention, the one or more polynucleotides of the invention or the one or more vectors of the invention.

The invention further provides a partially or fully differentiated cell descended from the cell or cell line of the invention, comprising the zinc finger nuclease of the invention, the one or more polynucleotides of the invention or the one or more vectors of the invention.

The invention provides a pharmaceutical composition comprising the zinc finger nuclease of the invention, the one or more polynucleotides of the invention, the one or more vectors of the invention, and/or the cell or cell line of the invention.

The invention also provides an in vitro method of modifying an endogenous BCL11A enhancer sequence in a cell, the method comprising administering the zinc finger nuclease of the invention or the one or more polynucleotides of the invention to the cell such that the endogenous BCL11A enhancer sequence is modified, optionally wherein: (i) the method further comprises introducing an exogenous sequence into the cell such that the exogenous sequence is inserted into the endogenous BCL11A enhancer sequence; or (ii) the modification comprises a deletion.

The invention further provides a pharmaceutical composition of the invention for use in a method of treating a hemoglobinopathy, such as a thalassemia (e.g. beta-thalassemia) or sickle cell disease, in a patient, the method comprising administering the pharmaceutical composition to the patient, optionally wherein the patient is human.

The invention provides an in vitro method of producing a genetically modified cell comprising a genomic modification within an endogenous BCL11A enhancer sequence, the method comprising the steps of: a) contacting a cell with the one or more polynucleotides of the invention; b) subjecting the cell to conditions conducive to expressing the fusion protein from the polynucleotide; and c) modifying the endogenous BCL11A enhancer sequence with the expressed fusion protein sufficient to produce the genetically modified cell, optionally wherein the method further comprises stimulating the cell with at least one cytokine.

The invention also provides a kit comprising the zinc finger nuclease of the invention, the one or more polynucleotides of the invention, the one or more vectors of the invention and/or the cell or cell line of the invention.

The invention further provides a cell or cell line of the invention for use in a method of treating or preventing a hemoglobinopathy, such as a thalassemia (e.g. beta-thalassemia) or sickle cell disease.

[0017] Disclosed herein is a non-naturally occurring zinc finger protein comprising a zinc finger protein (ZFP) comprising 4, 5 or 6 fingers, each finger comprising a recognition helix region that recognizes a DNA target subsite wherein the recognition helix regions comprise the sequences in the order shown in a single row of Table 1. Within each zinc finger, the 7 amino acid recognition helix region is numbered -1 to +6 within the zinc finger backbone (of approximately 30 residues, including zinc coordinating residues). 1, 2, 3 or more of the component zinc fingers of the zinc finger proteins described herein may further comprise mutations to one or more residues outside the recognition helix region, including but not limited to mutations to amino acids at position -5, position -14 or at both positions -5 and -14 (numbering continuing from the -1 to +6 numbering used for the recognition helix region) are mutated. See, e.g., Qm4 and Qm14 mutations described in U.S. Provisional Patent Applications 62/378,978 and 62/443,981. The component zinc fingers of the zinc finger protein can be linked by any linkers, for example as described in U.S. Patent No. 8,772,453. The ZFP comprises the recognition helixes as shown in Table 1 for the proteins designated as follows: 63014 (which binds to the target site shown in SEQ ID NO:1) and 65722 (which binds to the target site shown in SEQ ID NO:2).

[0018] In certain aspects of the disclosure, the zinc finger proteins as described herein may be fused to a functional domain (e.g., transcriptional activation domain, transcriptional repression domain, cleavage domain (to form a zinc finger nuclease), etc.). Any linker may be used to operably link the cleavage domain and the zinc finger protein, including but not limited to linkers as described in U.S. Patent Nos. 9,394,531 and 9,567,609. Furthermore, when a FokI cleavage domain is used, further mutations in the catalytic domain, dimerization domain, to phosphate contact residues (not in the dimerization or catalytic domain), and

combinations of mutations in any one of the catalytic domain, dimerization domain and to phosphate contact residues may be present, including but not limited to ELD or KKR mutations to the dimerization domain, mutations to residues 525 (K to S) of the FokI domain, and combinations of ELD or KKR mutations to the dimerization domain and mutations to residues 525 (K to S) of the FokI domain, numbered relative to wild-type. See, U.S. Patent Nos. 7,888,121; 7,914,796; 8,034,598; 8,623,618 and U.S. Patent Publication No. 2011/0201055 and U.S. Provisional Patent Application Nos. 62/378,978 and 62/443,981.

[0019] Zinc finger nucleases (ZFNs) may be used in dimerizing pairs to cleave at or near one or both of the target sites for the ZFNs of the pair, for example, the "left partner" of Table 1 (e.g., 63014) can form a dimer with the "right partner" of Table 1 (e.g., 65722) to cleave BCL11A enhancer sequences. For example, a pair of ZFNs may comprise the following amino acid sequences:

63014: MDYKDHDGDYKDHDIDYKDDDDKMAPKKRKVGIVHGVPAAMAERPFQCRI
 CMQNFSDQSNLRAHIRTHTGEKPFACDICGRKFARNFSLTMETKIHTGSQKPF
 QCRICMQNFSSGTNLNHIHTHTGEKPFACDICGRKFATSGSLTRHTKIHTHPR
 APIPKPFQCRICMQNFSDQSNLRAHIRTHTGEKPFACDICGRKFAAQCCFLHHT KIH-Linker -
 ELEEKSELRHKLKYVPHEYIELIEIARNSTQDRILEMKVMEFFMKVYGYRGK
 HLGGSRKPDGAIYTVGSPIDYGVIVDTKA YSGGYNLPIGQADEMERYVEENQ
 TRDKHLNPNEWWKVYPSSVTEFKFLFVSGHFGKNYKAQLTRLNHNITNCNGA
 VLSVEELLIGGEMIKAGTLTLEEVRRKFNNGEINFRS (SEQ ID NO:3); and
 MDYKDHDGDYKDHDIDYKDDDDKMAPKKRKVGIVHGVPAAMAERPFQCRI
 CMQKFARNDHRTTHTKIHTGEKPFQCRICMQNFSSQKAHLIRHIRTHTGEKPF
 CDICGRKFAQKGTLEHTKIHTGSQKPFQCRICMQNFSRGRDLSRHIRTHTGE
 KPFACDICGRKFARRDNLHSHTKIH-Linker-
 ELEEKSELRHKLKYVPHEYIELIEIARNSTQDRILEMKVMEFFMKVYGYRGK
 HLGGSRKPDGAIYTVGSPIDYGVIVDTKAYSGGYNLPIGQADEMQRYVKENQ
 TRNKHINPNEWWKVYPSSVTEFKFLFVSGHFGKNYKAQLTRLNRKTNCNGA
 VLSVEELLIGGEMIKAGTLTLEEVRRKFNNGEINF (SEQ ID NO:4), wherein the Linker sequence can be any

linker sequence known in the art, for example as described in U.S. Patent Nos. 9,394,531 and 9,567,609. The linker for 63014 may comprise or consist of the L7c5 linker (LRGSISRARPLNPHP (SEQ ID NO:5)) and the linker used in 65722 comprises or consists of the L0 linker (LRGSQLVKS (SEQ ID NO:6), see U.S. Patent No. 9,567,609). The FokI cleavage domain sequence C-terminal to the Linker of the sequences shown above may also comprise alternative FokI domains operably linked to the zinc finger protein. The FokI cleavage domain may include alternate or additional mutations to the catalytic domain, the dimerization domain, phosphate contact residues, and combinations of mutations to any one of the catalytic domain, dimerization domain and phosphate contact residues.

[0020] Described herein is the delivery of at least one nuclease (e.g., a nuclease that binds to a BCL11A enhancer sequence) to a human stem cell or precursor cell (HSC/PC) for the purpose of genome engineering. In certain aspects of the disclosure, the nuclease comprises a zinc finger protein (ZFP) comprising 4, 5 or 6 fingers, each finger comprising a recognition helix region that recognizes a target subsite wherein the recognition helix regions comprise the sequences in the order shown in a single row of Table 1. The ZFN nuclease of the invention comprises the pair of nucleases designated 63014/65722. The nuclease(s) as described herein may further comprise a linker (e.g., between the DNA-binding domain and the cleavage domain), for example a linker as shown in 9,567,609, including but not limited to LRGSISRARPLNPHP (SEQ ID NO:5) or (LRGSQLVKS (SEQ ID NO:6)).

[0021] In some embodiments, the nuclease is delivered as a peptide, while in others it is delivered as a nucleic acid encoding the at least one nuclease. In some embodiments, more than one nuclease is used. In some preferred embodiments, the nucleic acid encoding the nuclease is an mRNA, and in some instances, the mRNA is protected. In some aspects, the mRNA may be chemically modified (See e.g. Kormann et al. (2011) Nature Biotechnology 29(2): 154-157). In other aspects, the mRNA may comprise an ARCA cap (see U.S.

Patent Nos. 7,074,596 and 8,153,773). In further embodiments, the mRNA may comprise a mixture of unmodified and modified nucleotides (see U.S. Patent Publication No. 2012/0195936). In a preferred embodiment, the nucleic acid encoding the nuclease(s) is delivered to the HSC/PC via electroporation. In some embodiments, the nuclease cleaves at or near the binding site of a transcription factor. In some aspects, the transcription factor is GATA-1.

[0022] Disclosed herein is a cell or cell line in which an endogenous BCL11A enhancer sequence is genetically modified by a nuclease as described herein (e.g., shown in Table 1), for example as compared to the wild-type sequence of the cell. The genetic modification to the BCL11A enhancer results in modification of globin (beta and gamma) gene expression. Nuclease-modified cells or cell lines as described herein are distinguishable in structure, function, and combinations of both structure and function from wild-type. The genetically modified cell or cell lines may be heterozygous or homozygous for the modification. The modifications may comprise insertions (e.g., transgene insertion) deletions, and combinations of insertions and deletions; such insertions, deletions, and combinations of insertions and deletions are commonly referred to as "indels". In some preferred embodiments, indels result in the destruction of a transcription factor binding site. In certain embodiments, the modification is at or near the nuclease(s) binding site(s), cleavage site(s), and combinations of binding and cleavage sites, for example, within 1-300 (or any value therebetween) base pairs upstream or downstream of the site(s) of cleavage, more preferably within 1-100 base pairs (or any value therebetween) of either side of the binding site(s), cleavage site(s), and combinations of binding and cleavage sites shown in Table 1, even more preferably within 1 to 50 base pairs (or any value therebetween) on either side of the binding site(s), cleavage site(s), and combinations of binding and cleavage sites. In certain embodiments, the genetic modification of the BCL11A enhancer sequence is within and/or between sequences shown in Table 1 (target sites). The modification may also include modifications to one or more nucleotides in the cleavage sites. The modification may also include modifications to one or more nucleotides in the binding sites. The modification may further include modifications to one or more nucleotides in the cleavage sites, and in one or more of the binding sites. In certain embodiments, one or more of the nuclease target site(s) is(are) not modified. In other embodiments, at least one of the target sites for the nuclease(s) is(are) modified. In certain embodiments, the modification is at or near the "+58" region of the BCL11A enhancer, for example, at or near a nuclease binding site shown in any of SEQ ID NO: 1 and SEQ ID NO:2. Any cell or cell line may be modified by the nucleases as described herein, for example a stem cell (hematopoietic stem cell such as a CD34+ hematopoietic stem cell) or red blood cell (RBC) precursor cell.

[0023] Also described are cells or cell lines obtained following modification by a nuclease as described herein, for example cells or cell lines descended from a nuclease-modified cell or cell line as described herein. Partially or fully differentiated cells descended from the modified stem cells as described herein are also provided (e.g., RBCs or RBC precursor cells). The cells descended from the nuclease-modified cells may be propagated, differentiated, and combinations of both propagated and differentiated *in vitro* (culture) or may differentiate within a live subject, for example following *ex vivo* administration of a nuclease-modified stem cell. Any of the genetically modified cells or cell lines disclosed herein may show increased expression of gamma globin. Compositions such as pharmaceutical compositions comprising the genetically modified cells as described herein are also provided.

[0024] Delivery of a donor nucleic acid to a target cell may provide a genetically modified cell in which the donor is integrated into the cell. The donor may be delivered prior to, after, or along with the nucleic acid encoding the nuclease(s) of Table 1. The donor nucleic acid may comprise an exogenous sequence (transgene) to be integrated into the genome of the cell, for example, an endogenous locus. In some embodiments, the donor may comprise a full-length gene or fragment thereof flanked by regions of homology with the targeted cleavage site. In some embodiments, the donor lacks homologous regions and is integrated into a target locus through homology independent mechanism (i.e. NHEJ). The donor may comprise any nucleic acid sequence, for example a nucleic acid that, when used as a substrate for homology-directed repair of the nuclease-induced double-strand break, leads to a donor-specified deletion to be generated at the

endogenous chromosomal locus (e.g., BCL11A enhancer region) or, alternatively (or in addition to), novel allelic forms of (e.g., point mutations that ablate a transcription factor binding site) the endogenous locus to be created. In some aspects, the donor nucleic acid is an oligonucleotide wherein integration leads to a gene correction event, or a targeted deletion.

[0025] In other aspects, the nuclease, donor, and combinations of both the nuclease and donor is(are) delivered by viral, non-viral, and combinations of viral and non-viral gene transfer methods. In preferred embodiments, the donor is delivered to the cell via an adeno-associated virus (AAV). In some instances, the AAV comprises LTRs that are of a heterologous serotype in comparison with the capsid serotype.

[0026] In some aspects, deletions comprising regions within the DNaseI hypersensitive regions of the enhancer (e.g., the +58 region of the BCL11A enhancer) are made using one or more nucleases as shown in Table 1. These deletions can comprise from about 1 nucleotide to about 551 nucleotides. Thus, the deletions can comprise, 1, 5, 10, 15, 20, 25, 30, 40, 50, 100, 150, 200, 250, 300, 350, 400, 450, 500, 550 nucleotides, or any value therebetween. In some embodiments, the deletions comprise binding regions for one or more transcription factors. In some preferred embodiments, the deletions comprise a GATA-1 binding site, or the binding site for GATA-1 in combination with other factors.

[0027] In some aspects of the disclosure, the DNA binding domains of Table 1 are fused to a functional domain. Some aspects include fusion of the DNA binding domains with domains capable of regulating the expression of a gene. In some aspects of the disclosure, the fusion proteins comprise the DNA binding domain of Table 1 fused to a gene expression modulatory domain where the modulator represses gene expression.

[0028] In some aspects of the disclosure, the HSC/PC cells are contacted with the nucleases, the DNA binding proteins, and combinations of the nucleases and DNA binding proteins of the disclosure (*i.e.*, ZFPs as shown in Table 1). The nucleases, the DNA binding proteins, and combinations of the nucleases and DNA binding proteins may be delivered as nucleic acids or as proteins. The nucleic acids may be mRNAs encoding the DNA binding proteins, and combinations of the nucleases and DNA binding proteins, and in further instances, the mRNAs may be protected. The mRNA may be chemically modified, may comprise an ARCA cap, a mixture of unmodified and modified nucleotides, and combinations of an ARCA cap and a mixture of unmodified and modified nucleotides. Cells or cell lines descended from these cells are also provided, including partially or fully differentiated cells.

[0029] In some aspects, the HSC/PC are contacted with the nucleases, the DNA binding proteins, and combinations of the nucleases and DNA binding proteins of the disclosure *ex vivo*, following apheresis of the HSC/PC from a subject, or purification from harvested bone marrow. In some embodiments, the nucleases described herein cause modifications within the BCL11A enhancer regions, for example resulting a genetically modified cell that is structurally, functionally, and combinations of structurally and functionally distinct from wild-type cells, other modified (e.g., nuclease-modified) cells, and combinations of wild-type and other modified cells. In further embodiments, the HSC/PC containing the BCL11A enhancer region modifications are introduced back into the subject. In some instances, the HSC/PC containing the BCL11A enhancer region modifications are expanded prior to introduction. In other aspects, the genetically modified HSC/PCs are given to the subject in a bone marrow transplant wherein the HSC/PC engraft, differentiate and mature *in vivo*. In some embodiments, the HSC/PC are isolated from the subject following G-CSF-induced mobilization, plerixafor-induced mobilization, and combinations of G-CSF- and plerixafor-induced mobilization, and in others, the cells are isolated from human bone marrow or human umbilical cords. In some aspects, the subject is treated to a mild myeloablative procedure prior to introduction of the graft comprising the modified HSC/PC, while in other aspects, the subject is treated with a vigorous myeloablative conditioning regimen. In some embodiments, the compositions of the invention are used to treat or prevent a hemoglobinopathy. In some aspects, the hemoglobinopathy is a thalassemia. In some aspects, the hemoglobinopathy is a beta thalassemia, while in other aspects, the hemoglobinopathy is sickle cell disease.

[0030] In some embodiments, the HSC/PC are further contacted with a donor molecule. In some embodiments, the donor molecule is delivered by a viral vector. The donor molecule may comprise one or more sequences encoding a functional polypeptide (e.g., a cDNA or fragment thereof), with or without a promoter. Additional sequences (coding or non-coding sequences) may be included when a donor molecule is used for inactivation, including but not limited to, sequences encoding a 2A peptide, SA site, IRES, *etc.*

[0031] In one aspect, the HSC/PC are contacted by the compositions of the invention *in vivo*. The nucleases, DNA binding proteins, or combination of nucleases and DNA binding proteins may be delivered to HSC/PC *in situ* by methods known in the art. In some embodiments, the nucleases of the invention comprise a viral particle that is administered to the subject in need, while in others, the nucleases, DNA binding proteins, or combination of nucleases and DNA binding proteins comprise a nanoparticle (e.g. liposome). In some embodiments, the viral particles, nanoparticles, or combination of viral particles and nanoparticles are delivered to the organ (e.g. bone marrow) wherein the HSC/PC reside.

[0032] In another aspect, described herein are methods of integrating a donor nucleic acid into the genome of a cell via homology-independent mechanisms. The methods comprise creating a double-stranded break (DSB) in the genome of a cell and cleaving the donor molecule using a nuclease as described herein, such that the donor nucleic acid is integrated at the site of the DSB. In certain embodiments, the donor nucleic acid is integrated via non-homology dependent methods (e.g., NHEJ). As noted above, upon *in vivo* cleavage the donor sequences can be integrated in a targeted manner into the genome of a cell at the location of a DSB. The donor sequence can include one or more of the same target sites for one or more of the nucleases used to create the DSB. Thus, the donor sequence may be cleaved by one or more of the same nucleases used to cleave the endogenous gene into which integration is desired. In certain embodiments, the donor sequence includes different nuclease target sites from the nucleases used to induce the DSB. DSBs in the genome of the target cell may be created by any mechanism. In certain embodiments, the DSB is created by one or more zinc-finger nucleases (ZFNs), fusion proteins comprising a zinc finger binding domain that is engineered to bind a sequence within the region of interest, and a cleavage domain or a cleavage half-domain.

[0033] In one aspect, the donor may encode a regulatory protein of interest (e.g. ZFP TFs, TALE TFs or a CRISPR/Cas TF) that binds to, modulates expression of, or both binds to and modulates expression of a gene of interest. In one embodiment, the regulatory proteins bind to a DNA sequence and prevents binding of other regulatory factors. In another embodiment, the binding of the regulatory protein may modulate (*i.e.* induce or repress) expression of a target DNA.

[0034] In some embodiments, the transgenic HSC/PC cell, transgenic animal, or combination of transgenic HSC/PC cell and animal includes a transgene that encodes a human gene. In some instances, the transgenic animal comprises a knock out at the endogenous locus, and replacement of the endogenous gene with its human counterpart, thereby allowing the development of an *in vivo* system where the human protein may be studied in isolation. Such transgenic models may be used for screening purposes to identify small molecules or large biomolecules or other entities which may interact with or modify the human protein of interest. In some aspects, the transgene is integrated into the selected locus (e.g., safe-harbor) into a stem cell (e.g., an embryonic stem cell, an induced pluripotent stem cell, a hematopoietic stem cell, *etc.*) or non-human animal embryo obtained by any of the methods described herein, and then the embryo is implanted such that a live non-human animal is born. The animal is then raised to sexual maturity and allowed to produce offspring wherein at least some of the offspring comprise edited endogenous gene sequence or the integrated transgene.

[0035] In another aspect, provided herein is a method of altering gene expression (e.g., BCL11A, a globin gene, and combinations of BCL11A and a globin gene) in a cell, the method comprising: introducing, into the cell, one or more nucleases as described herein (shown in Table 1), under conditions such that the one or more proteins are expressed and expression of the gene is altered. In certain embodiments, expression of a

globin gene (e.g., gamma globin or beta globin) is altered (e.g., increased). Any of the methods described herein may further comprise integrating a donor sequence (e.g., transgene or fragment thereof under the control of an exogenous or endogenous promoter) into the genome of the cell, for example integrating a donor at or near the site of nuclease cleavage in the BCL11A gene. The donor sequence is introduced to the cell using a viral vector, as an oligonucleotide, on a plasmid, and combinations of one or more methods selected from using a viral vector, as an oligonucleotide, or on a plasmid. The cell in which gene expression is altered may be, for example, a red blood cell (RBC) precursor cell, a hematopoietic stem cell (e.g., CD34+ cell), and combinations of RBC precursor cell and a hematopoietic stem cell.

[0036] Described herein is a method of producing a genetically modified cell comprising a genomic modification within an endogenous BCL11A enhancer sequence (a modification to the nucleotide sequence of the BCL11A enhancer sequence), the method comprising the steps of: a) contacting a cell with a polynucleotide (e.g. DNA or mRNA) encoding a zinc finger nuclease comprising 4, 5, or 6 zinc finger domains in which each of the zinc finger domains comprises a recognition helix region in the order shown in a single row of Table 1; b) subjecting the cell to conditions conducive to expressing the zinc finger protein from the polynucleotide; and c) modifying the endogenous BCL11A enhancer sequence with the expressed zinc finger protein sufficient to produce the genetically modified cell. The cells may be stimulated with at least one cytokine (e.g., prior to step (a)). The polynucleotide may be contacted with the cell using any suitable method, including but not limited, via transfection, using a non-viral vector, using a viral vector, by chemical means or by exposure to an electric field (e.g., electroporation).

[0037] Cells comprising one or a combination of the genomic modifications described herein are also provided, including cells descended from the cells produced by the methods described herein.

[0038] Also disclosed is a method of treating a patient in need of an increase in globin gene expression, the method comprising administering to the patient the pharmaceutical preparation, wherein the pharmaceutical preparation comprises genetically modified cells, proteins, polynucleotides, and combinations of one or more selected from genetically modified cells, proteins, and polynucleotides, as described herein in an amount sufficient to increase the globin gene expression in the patient. The patient may be known to have, is suspected of having, or is at risk of developing a thalassemia or sickle cell disease.

[0039] A kit, comprising the nucleic acids, proteins, genetically modified cells, and combinations of one or more selected from the nucleic acids, proteins, and genetically modified cells of the invention, is also provided. The kit may comprise nucleic acids encoding the nucleases, (e.g. RNA molecules or ZFN, TALEN or CRISPR/Cas system encoding genes contained in a suitable expression vector), aliquots of the nuclease proteins, donor molecules, suitable modifiers of stem cell self-renewal ("stemness"), cells, buffers, instructions (e.g., for performing the methods of the invention) and the like, including various combinations of these kit components. Disclosed is a genetically modified cell (e.g., stem cell such as a hematopoietic (CD34+) stem cell or RBC precursor cell) comprising at least one genomic modification made by a nuclease (e.g., as shown in a single row of Table 1), wherein the genomic modification is within an endogenous BCL11A enhancer sequence, and further wherein the genomic modification is selected from the group consisting of insertions, deletions and combinations thereof and comprises a modification at, near or between any of SEQ ID NO: 1 and SEQ ID NO:2. In certain embodiments, the cell is a genetically modified differentiated cell descended from a stem cell as described herein (e.g., a RBC descended from a hematopoietic stem cell or RBC precursor cell).

[0040] The nuclease may comprise at least one zinc finger nuclease (ZFN) (e.g., as shown in Table 1), at least one TALEN, and combinations of at least one ZFN and at least one TALEN. The nuclease(s) may be introduced into the cell in protein form, as a polynucleotide encoding the nuclease(s), or as a combination of protein form and polynucleotide encoding the nuclease(s). In certain embodiments, the genomic modification comprises an insertion that comprises integration of a donor polynucleotide encoding a transgene. Also

described are pharmaceutical compositions comprising one or more of the genetically modified cells as described herein.

[0041] Also described is a DNA-binding protein comprising a zinc finger protein comprising 4, 5 or 6 zinc finger domains comprising a recognition helix region, wherein the zinc finger proteins comprise the recognition helix regions in the order shown in a single row of Table 1. Also described is a TALE protein comprising a plurality of repeats that bind to a sequence comprising a portion (e.g., at least 4, 5, 6 or more) base pairs of the target sites shown in Table 1. A fusion protein comprising a zinc finger protein or TALE protein as described herein and a wild-type or engineered cleavage domain or cleavage half-domain is also described as are polynucleotides encoding the proteins (ZFPs, TALEs, ZFNs, TALENs) as described herein. Cells (e.g., isolated stem cells such as hematopoietic (CD34+) stem cells) comprising one or more polynucleotides, proteins, and combinations of polynucleotides and proteins as described herein are also described. Also described are kits comprising one or more proteins, polynucleotides, cells, or combinations thereof as described herein.

[0042] A method of altering globin gene expression in a cell (e.g., RBC precursor cell, hematopoietic stem cell and combinations of RBC precursor cell and hematopoietic stem cell) is also described, the method comprising: introducing, into the cell, one or more polynucleotides encoding one or more nucleases as described herein, under conditions such that the one or more proteins are expressed and expression of the globin gene (e.g., gamma globin, beta globin, and combinations of gamma and beta globin) is altered (e.g., increased). The methods may further comprise integrating a donor sequence into the genome of the cell, for example using a viral vector, as an oligonucleotide or on a plasmid. The donor sequence may comprise a transgene under the control of an endogenous or exogenous promoter.

[0043] Also described is a method of producing a genetically modified cell comprising a genomic modification within an endogenous BCL11A enhancer sequence (e.g., target site as shown in Table 1), the method comprising the steps of: (a) contacting a cell with a polynucleotide encoding a fusion protein comprising a zinc finger nuclease comprising 4, 5, or 6 zinc finger domains in which each of the zinc finger domains comprises a recognition helix region in the order shown in a single row of Table 1; (b) subjecting the cell to conditions conducive to expressing the fusion protein from the polynucleotide; and (c) modifying the endogenous BCL11A enhancer sequence with the expressed fusion protein sufficient to produce the genetically modified cell. The method may further comprise stimulating the cells with at least one cytokine. The polynucleotide(s) may be delivered inside the cell, for example using a non-viral delivery system, a viral delivery system, a delivery vehicle, and combinations selected from a non-viral delivery system, a viral delivery system, and a delivery vehicle and may comprise subjecting the cells to an electric field or employing cell membrane disruption as a delivery mechanism (so called 'Squeeze Technology', see e.g. Sharei et al. (2015) PLOS ONE doi: 10.1371/journal.pone.0118803).

[0044] Methods of treating a patient in need of an increase in globin gene expression (e.g., a patient is known to have, is suspected of having, or is at risk of developing a hemoglobinopathy such as a thalassemia (e.g., β -thalassemia) or sickle cell disease are also described, the method comprising administering to the patient the pharmaceutical composition as described herein (e.g., proteins, polynucleotides, cells or a combination selected from proteins, polynucleotides and cells) in an amount sufficient to increase the globin gene expression in the patient.

[0045] These and other aspects will be readily apparent to the skilled artisan in light of disclosure as a whole.

BRIEF DESCRIPTION OF THE DRAWINGS

[0046]

Figures 1A and 1B are schematics depicting an overview of the oligonucleotide duplex integration site assay. Figure 1 shows the first step of the assay, in which cells are treated with ZFNs in the presence of supplemental oligonucleotide duplex DNA, which is captured into a fraction of the resulting cleavage events. Figure 1B shows the subsequent step in which cells are cultured for seven days, genomic DNA is isolated, and segments of the genome flanking donor integration sites are amplified via adaptor-mediated PCR using a primer to the integrated oligonucleotide duplex. Amplicons are then sequenced to reveal candidate cleavage sites.

Figure 2 depicts the location of 455 potential off-target loci identified by the oligonucleotide duplex integration site assay using the 51857/51949 original or parent ZFN pair. The top 63 loci are highlighted in gray and were analyzed in follow up indel analyses. Each locus is identified by the chromosome and base number indicating the most likely location of cleavage, as well as an indication of the number of integrants detected at that locus.

Figures 3A through 3C depict exemplary phosphate contacting residues within the zinc finger scaffold. Figure 3A depicts a zinc finger interaction with a DNA molecule, and shows the location of the wild type arginine side chain and how it interacts with the phosphate backbone of the DNA molecule. Figure 3B (SEQ ID Nos:7 to 17) depicts exemplary ZFN sequences showing where this arginine is located in the primary sequence of each zinc finger (vertical row of 'R' residues indicated via bold arrow) and also highlights those arginines that were substituted with glutamine in the ZFP backbone to eliminate the corresponding phosphate contact (individually boxed 'R' residues). The sequences also show the recognition helix regions (where residues -1 to +3, +5 and +6 are boxed and shaded) as well as a portion of the linker between the C-terminal zinc finger domain and the cleavage domain (cleavage domain not shown). Figure 3C further depicts the spatial location of another potential backbone contacting lysine sidechain present in the FokI cleavage domain, which during specificity optimization may be substituted to a serine.

Figure 4 is a graph comparing modification levels via treatment of CD34 cells with either the original 51857/51949 pair or with the optimized 63014/65722 pair at various tested loci. Shown across the bottom or on the x-axis, are the loci identified as potential cleavage targets, with the percent indels for each site shown on the vertical or y-axis. Note that the y-axis is shown in a log scale. The dark gray bars show the loci cleaved by the 51857/51949 pair, and the amount of cleavage detected, while the light gray bars are those loci cleaved by the optimized pair where nearly all cleavage measured is at the targeted BCL11a target sequence.

Figures 5A and 5B are graphs depicting the relative ratios of globin mRNAs made in hCD34+ cells following treatment with BCL11A specific ZFNs and erythroid differentiation. CD34+ cells derived from two healthy human donors (PB-MR-003 or PB-MR-004) were treated or not treated with the ZFN pair and then α , β , and γ globin expression was analyzed. The best method to determine the amount of γ globin mRNA found following ZFN treatment is to express the change in expression as either a ratio of γ globin to β globin (Figure 5A), or γ globin to α globin mRNA (Figure 5B).

Figure 6 is a graph depicting the relative amounts of γ -globin protein produced in the treated CD34+ cells. As above, two CD34+ cell lots derived from healthy human donors were used (PB-MR-003 and PB-MR-004). In this experiment, an approximate 3-4 fold elevation of fetal globin protein percentages to levels of about 15%-20% was observed in erythroid progeny of HSPCs upon 63014/65722-mediated disruption of the BCL11A enhancer in both donor lots.

Figure 7 is a graph indicating relative human chimerism in mice engrafted with 63014/65722 treated donor lots ("ZFN") as described above. Human chimerism was measured through the detection of cells bearing a hCD45 marker on their surface using FACS. Percentages of human hCD45+ cells in the peripheral blood collected at either 8 or 12 weeks post-transplant are indicated. The data showed good engraftment levels in this study with comparable human chimerism following engraftment of untransfected control ("(-)") and ZFN transfected HSPC ("ZFN"). Open circles and triangles represent individual animals.

Figure 8 shows the percentage of chimerism detected in the bone marrow of the engrafted mice where human cells were identified by the presence of hCD45 on their cell surfaces. Samples were analyzed at 12

weeks post-engraftment.

Figures 9A through 9D are graphs depicting the reconstitution of various hematopoietic cell lineages tested by FACS analysis of bone marrow cells in engrafted mice obtained at week 12 with antibodies recognizing lineage specific cell surface markers. The data showed comparable representation of all analyzed human hematopoietic lineages in the bone marrow at week 12 post-injection between the BCL11A-specific ZFN mRNA treated CD34+ cell progeny ("14/22") and that of the untransfected cells ("(-)"). Shown are data from lymphoid, myeloid, erythroid and HSPC (Figures 9A through 9D, respectively) for cells derived from both donors ("003" and "004"). The data showed comparable representation of all analyzed human hematopoietic lineages in the bone marrow at week 12 post-injection between the Bcl11A ZFN mRNA treated CD34+ cell progeny and that of the untransfected cells.

Figure 10 is a graph depicting the level of gene modification at the BCL11A target in DNA isolated from the peripheral blood of engrafted mice, assayed by deep sequencing. Data are shown for the input cells (2 days after ZFN transfection, ("+")), and then for blood cells 8 or 12 weeks following engraftment, and demonstrated a good retention of gene modification. Untransfected cells are represented by "(-)" in the Treatment line.

Figure 11 is a graph depicting the amount of gene modification at the BCL11A target for bone marrow cell samples following engraftment of the ZFN-treated cells ("+"). Untreated cells are represented by "(-)" in the Treatment line. Comparable modification was observed in both BCL11A dependent lineages (B cells, expressing the CD19 marker; primitive progenitors, expressing CD45 and high levels of CD38) and BCL11A independent (myeloid) lineages. Although the input gene modification levels were higher in the PB-MR-003 donor sample than in the PB-MR-004 donor sample, the PB-MR-004 derived cells consistently showed higher modification levels, *i.e.* better retention of modification, in mice than those derived from PB-MR-003.

Figure 12 is a graph depicting the amount of gene modification in erythroid cells derived from week 12 bone marrow cells following *in vitro* differentiation for 14 days. The data were from mice originally engrafted with the two different donors described above, and demonstrated that the BCL11A modification mediated by ZFN treatment ("+" ZFN) is not markedly altered during the erythroid differentiation. Cells that were not treated with ZFN are indicated by "(-) ZFN".

Figures 13A and 13B are graphs depicting the relative amount of γ -globin encoding mRNA, where the concentration of γ -globin mRNA is depicted either as a ratio of γ globin/ β globin mRNA (Figure 9A) or as a ratio of γ globin / α globin mRNA (Figure 9B). Both in the untransfected ("(-) ZFN") and the ZFN treated samples ("+" ZFN") γ -globin to β globin or γ -globin to α -globin mRNA ratios differed widely between the erythroid progenies of individual mice from the same group. However, despite this variability and the variability introduced by the use of two different human donors, the 63014/65722 treated sample averages show an ~1.5-2 fold increase in γ globin mRNA levels compared to their respective untransfected counterparts.

Figure 14 is a graph depicting the difference in the amount of γ -globin protein (expressed as either a ratio of γ -globin/ α -globin or γ -globin/total β -like protein) in bone marrow derived cells from engrafted mice where the bone marrow cells were submitted to an *in vitro* differentiation protocol. Protein levels were measured 16 days into differentiation. The Gamma (γ) globin (sum of the Agamma and Ggamma peaks) to alpha (α) globin ratios were determined, as well as the Gamma globin (sum of the Agamma and Ggamma peaks)/ over beta-like globin ratios (sum of the Agamma, Ggamma, beta and delta- globin peaks). In line with the poor erythroid differentiation of the PB-MR-003 derived samples the gamma-globin levels in the untransfected cells derived from this donor were very high (~30%), and therefore ZFN treatment ("+" ZFN") resulted in only a 1.2-fold increase in gamma-globin levels as compared with untreated cells ("(-) ZFN"). The PB-MR-004 showed more typical untransfected levels (~9%) and exhibited an ~2-fold increase in gamma-globin protein levels after 12 weeks passage through the mouse.

DETAILED DESCRIPTION

[0047] Disclosed herein are compositions and methods for genome engineering for the modulation of BCL11A, gamma globin, and combinations of BCL11A and gamma globin expression and for the treatment, prevention, or treatment and prevention of hemoglobinopathies. In particular, via targeting with nucleases comprising the ZFPs having the recognition helix regions as shown in a single row of Table 1, disruption of an enhancer of BCL11A is efficiently achieved in HSC/PC and results in a change in relative gamma globin expression during subsequent erythropoiesis. This modulation of BCL11A and gamma globin expression is particularly useful for treatment of hemoglobinopathies (e.g., beta thalassemias, sickle cell disease) wherein there is insufficient beta globin expression or expression of a mutated form of beta-globin. Using the compositions of the invention, the complications and disease related sequelae caused by the aberrant beta globin can be overcome by alteration of the expression of gamma globin in erythrocyte precursor cells.

General

[0048] Practice of the methods, as well as preparation and use of the compositions disclosed herein employ, unless otherwise indicated, conventional techniques in molecular biology, biochemistry, chromatin structure and analysis, computational chemistry, cell culture, recombinant DNA and related fields as are within the skill of the art. These techniques are fully explained in the literature. See, for example, Sambrook et al. MOLECULAR CLONING: A LABORATORY MANUAL, Second edition, Cold Spring Harbor Laboratory Press, 1989 and Third edition, 2001; Ausubel et al., CURRENT PROTOCOLS IN MOLECULAR BIOLOGY, John Wiley & Sons, New York, 1987 and periodic updates; the series METHODS IN ENZYMOLOGY, Academic Press, San Diego; Wolffe, CHROMATIN STRUCTURE AND FUNCTION, Third edition, Academic Press, San Diego, 1998; METHODS IN ENZYMOLOGY, Vol. 304, "Chromatin" (P.M. Wassarman and A. P. Wolffe, eds.), Academic Press, San Diego, 1999; and METHODS IN MOLECULAR BIOLOGY, Vol. 119, "Chromatin Protocols" (P.B. Becker, ed.) Humana Press, Totowa, 1999.

Definitions

[0049] The terms "nucleic acid," "polynucleotide," and "oligonucleotide" are used interchangeably and refer to a deoxyribonucleotide or ribonucleotide polymer, in linear or circular conformation, and in either single- or double-stranded form. For the purposes of the present disclosure, these terms are not to be construed as limiting with respect to the length of a polymer. The terms can encompass known analogues of natural nucleotides, as well as nucleotides that are modified in the base, sugar, phosphate moieties (e.g., phosphorothioate backbones), and combinations selected from base, sugar and phosphate moieties. In general, an analogue of a particular nucleotide has the same base-pairing specificity; *i.e.*, an analogue of A will base-pair with T.

[0050] The terms "polypeptide," "peptide" and "protein" are used interchangeably to refer to a polymer of amino acid residues. The term also applies to amino acid polymers in which one or more amino acids are chemical analogues or modified derivatives of a corresponding naturally-occurring amino acids.

[0051] "Binding" refers to a sequence-specific, non-covalent interaction between macromolecules (e.g., between a protein and a nucleic acid). Not all components of a binding interaction need be sequence-specific (e.g., contacts with phosphate residues in a DNA backbone), as long as the interaction as a whole is sequence-specific. Such interactions are generally characterized by a dissociation constant (K_d) of 10^{-6} M^{-1} or lower. "Affinity" refers to the strength of binding: increased binding affinity being correlated with a lower K_d .

[0052] A "binding protein" is a protein that is able to bind to another molecule. A binding protein can bind to, for example, a DNA molecule (a DNA-binding protein), an RNA molecule (an RNA-binding protein), a protein molecule (a protein-binding protein), or can bind to a combination of molecules selected from a DNA molecule, an RNA molecule or a protein. In the case of a protein-binding protein, it can bind to itself (to form homodimers, homotrimers, *etc.*), it can bind to one or more molecules of a different protein or proteins, or it can bind to both itself and one or molecules of a different protein or proteins. A binding protein can have more than one type of binding activity. For example, zinc finger proteins have DNA-binding, RNA-binding and protein-binding activity.

[0053] A "zinc finger DNA binding protein" (or binding domain) is a protein, or a domain within a larger protein, that binds DNA in a sequence-specific manner through one or more zinc fingers, which are regions of amino acid sequence within the binding domain whose structure is stabilized through coordination of a zinc ion. The term zinc finger DNA binding protein is often abbreviated as zinc finger protein or ZFP.

[0054] A "TALE DNA binding domain" or "TALE" is a polypeptide comprising one or more TALE repeat domains/units. The repeat domains are involved in binding of the TALE to its cognate target DNA sequence. A single "repeat unit" (also referred to as a "repeat") is typically 33-35 amino acids in length and exhibits at least some sequence homology with other TALE repeat sequences within a naturally occurring TALE protein.

[0055] Zinc finger and TALE binding domains can be "engineered" to bind to a predetermined nucleotide sequence, for example via engineering (altering one or more amino acids) of the recognition helix region of a naturally occurring zinc finger or TALE protein. Therefore, engineered DNA binding proteins (zinc fingers or TALEs) are proteins that are non-naturally occurring. Non-limiting examples of methods for engineering DNA-binding proteins are design and selection. A designed DNA binding protein is a protein not occurring in nature whose design/composition results principally from rational criteria. Rational criteria for design include application of substitution rules and computerized algorithms for processing information in a database storing information of existing ZFP and/or TALE designs and binding data. See, for example, U.S. Patent Nos. 6,140,081; 6,453,242; 6,534,261 and 8,585,526; see also PCT Publication Nos. WO 98/53058; WO 98/53059; WO 98/53060; WO 02/016536 and WO 03/016496.

[0056] A "selected" zinc finger protein or TALE is a protein not found in nature whose production results primarily from an empirical process such as phage display, interaction trap or hybrid selection. See *e.g.*, U.S. Patent Nos. 5,789,538; 5,925,523; 6,007,988; 6,013,453; 6,200,759; 8,586,526; PCT Publication Nos. WO 95/19431; WO 96/06166; WO 98/53057; WO 98/54311; WO 00/27878; WO 01/60970 WO 01/88197, WO 02/099084.

[0057] "TtAgo" is a prokaryotic Argonaute protein thought to be involved in gene silencing. TtAgo is derived from the bacteria *Thermus thermophilus*. See, *e.g.*, Swarts *et al. ibid*, G. Sheng *et al.* (2013) *Proc. Natl. Acad. Sci. U.S.A.* 111, 652). A "TtAgo system" is all the components required including, for example, guide DNAs for cleavage by a TtAgo enzyme.

[0058] "Recombination" refers to a process of exchange of genetic information between two polynucleotides, including but not limited to, donor capture by non-homologous end joining (NHEJ) and homologous recombination. For the purposes of this disclosure, "homologous recombination (HR)" refers to the specialized form of such exchange that takes place, for example, during repair of double-strand breaks in cells via homology-directed repair mechanisms. This process requires nucleotide sequence homology, uses a "donor" molecule to template repair of a "target" molecule (*i.e.*, the one that experienced the double-strand break), and is variously known as "non-crossover gene conversion" or "short tract gene conversion," because it leads to the transfer of genetic information from the donor to the target. Without wishing to be bound by any particular theory, such transfer can involve mismatch correction of heteroduplex DNA that forms between the broken target and the donor, "synthesis-dependent strand annealing," in which the donor is used to

resynthesize genetic information that will become part of the target, related processes, or combinations thereof. Such specialized HR often results in an alteration of the sequence of the target molecule such that part or all of the sequence of the donor polynucleotide is incorporated into the target polynucleotide.

[0059] In the methods of the disclosure, one or more targeted nucleases as described herein create a double-stranded break (DSB) in the target sequence (e.g., cellular chromatin) at a predetermined site. The DSB may result in indels by homology-directed repair or by non-homology-directed repair mechanisms. Deletions may include any number of base pairs. Similarly, insertions may include any number of base pairs including, for example, integration of a "donor" polynucleotide, optionally having homology to the nucleotide sequence in the region of the break. The donor sequence may be physically integrated or, alternatively, the donor polynucleotide is used as a template for repair of the break via homologous recombination, resulting in the introduction of all or part of the nucleotide sequence as in the donor into the cellular chromatin. Thus, a first sequence in cellular chromatin can be altered and, in certain embodiments, can be converted into a sequence present in a donor polynucleotide. Thus, the use of the terms "replace" or "replacement" can be understood to represent replacement of one nucleotide sequence by another, (*i.e.*, replacement of a sequence in the informational sense), and does not necessarily require physical or chemical replacement of one polynucleotide by another.

[0060] In any of the methods described herein, additional pairs of zinc-finger proteins or TALEN can be used for additional double-stranded cleavage of additional target sites within the cell.

[0061] Any of the methods described herein can be used for insertion of a donor of any size, or partial or complete inactivation of one or more target sequences in a cell by targeted integration of donor sequence that disrupts expression of the gene(s) of interest. Cell lines with partially or completely inactivated genes are also described.

[0062] In any of the methods described herein, the exogenous nucleotide sequence (the "donor sequence" or "transgene") can contain sequences that are homologous, but not identical, to genomic sequences in the region of interest, thereby stimulating homologous recombination to insert a non-identical sequence in the region of interest. Thus, in certain embodiments, portions of the donor sequence that are homologous to sequences in the region of interest exhibit between about 80 to 99% (or any integer therebetween) sequence identity to the genomic sequence that is replaced. In other embodiments, the homology between the donor and genomic sequence is higher than 99%, for example if only 1 nucleotide differs as between donor and genomic sequences of over 100 contiguous base pairs. In certain cases, a non-homologous portion of the donor sequence can contain sequences not present in the region of interest, such that new sequences are introduced into the region of interest. In these instances, the non-homologous sequence is generally flanked by sequences of 50-1,000 base pairs (or any integral value therebetween) or any number of base pairs greater than 1,000, that are homologous or identical to sequences in the region of interest. In other embodiments, the donor sequence is non-homologous to the first sequence, and is inserted into the genome by non-homologous recombination mechanisms.

[0063] "Cleavage" refers to the breakage of the covalent backbone of a DNA molecule. Cleavage can be initiated by a variety of methods including, but not limited to, enzymatic or chemical hydrolysis of a phosphodiester bond. Both single-stranded cleavage and double-stranded cleavage are possible, and double-stranded cleavage can occur as a result of two distinct single-stranded cleavage events. DNA cleavage can result in the production of either blunt ends or staggered ends. In certain embodiments, fusion polypeptides are used for targeted double-stranded DNA cleavage.

[0064] A "cleavage half-domain" is a polypeptide sequence which, in conjunction with a second polypeptide (either identical or different) forms a complex having cleavage activity (preferably double-strand cleavage activity). The terms "first and second cleavage half-domains," "+ and - cleavage half-domains" and "right and

left cleavage half-domains" are used interchangeably to refer to pairs of cleavage half-domains that dimerize.

[0065] An "engineered cleavage half-domain" is a cleavage half-domain that has been modified so as to form obligate heterodimers with another cleavage half-domain (e.g., another engineered cleavage half-domain). See, *also*, U.S. Patent Nos. 7,888,121; 7,914,796; 8,034,598; 8,623,618 and U.S. Patent Publication No. 2011/0201055.

[0066] The term "sequence" refers to a nucleotide sequence of any length, which can be DNA or RNA; can be linear, circular or branched and can be either single-stranded or double stranded. The term "donor sequence" refers to a nucleotide sequence that is inserted into a genome. A donor sequence can be of any length, for example between 2 and 100,000,000 nucleotides in length (or any integer value therebetween or thereabove), preferably between about 100 and 100,000 nucleotides in length (or any integer therebetween), more preferably between about 2000 and 20,000 nucleotides in length (or any value therebetween) and even more preferable, between about 5 and 15 kb (or any value therebetween).

[0067] "Chromatin" is the nucleoprotein structure comprising the cellular genome. Cellular chromatin comprises nucleic acid, primarily DNA, and protein, including histones and non-histone chromosomal proteins. The majority of eukaryotic cellular chromatin exists in the form of nucleosomes, wherein a nucleosome core comprises approximately 150 base pairs of DNA associated with an octamer comprising two each of histones H2A, H2B, H3 and H4; and linker DNA (of variable length depending on the organism) extends between nucleosome cores. A molecule of histone H1 is generally associated with the linker DNA. For the purposes of the present disclosure, the term "chromatin" is meant to encompass all types of cellular nucleoprotein, both prokaryotic and eukaryotic. Cellular chromatin includes both chromosomal and episomal chromatin.

[0068] A "chromosome," is a chromatin complex comprising all or a portion of the genome of a cell. The genome of a cell is often characterized by its karyotype, which is the collection of all the chromosomes that comprise the genome of the cell. The genome of a cell can comprise one or more chromosomes.

[0069] An "episome" is a replicating nucleic acid, nucleoprotein complex or other structure comprising a nucleic acid that is not part of the chromosomal karyotype of a cell. Examples of episomes include plasmids and certain viral genomes.

[0070] An "accessible region" is a site in cellular chromatin in which a target site present in the nucleic acid can be bound by an exogenous molecule which recognizes the target site. Without wishing to be bound by any particular theory, it is believed that an accessible region is one that is not packaged into a nucleosomal structure. The distinct structure of an accessible region can often be detected by its sensitivity to chemical and enzymatic probes, for example, nucleases.

[0071] A "target site" or "target sequence" is a nucleic acid sequence that defines a portion of a nucleic acid to which a binding molecule will bind, provided sufficient conditions for binding exist.

[0072] An "exogenous" molecule is a molecule that is not normally present in a cell, but can be introduced into a cell by one or more genetic, biochemical or other methods. "Normal presence in the cell" is determined with respect to the particular developmental stage and environmental conditions of the cell. Thus, for example, a molecule that is present only during embryonic development of muscle is an exogenous molecule with respect to an adult muscle cell. Similarly, a molecule induced by heat shock is an exogenous molecule with respect to a non-heat-shocked cell. An exogenous molecule can comprise, for example, a functioning version of a malfunctioning endogenous molecule or a malfunctioning version of a normally-functioning endogenous molecule.

[0073] An exogenous molecule can be, among other things, a small molecule, such as is generated by a

combinatorial chemistry process, or a macromolecule such as a protein, nucleic acid, carbohydrate, lipid, glycoprotein, lipoprotein, polysaccharide, any modified derivative of the above molecules, or any complex comprising one or more of the above molecules. Nucleic acids include DNA and RNA, can be single- or double-stranded; can be linear, branched or circular; and can be of any length. Nucleic acids include those capable of forming duplexes, as well as triplex-forming nucleic acids. See, for example, U.S. Patent Nos. 5,176,996 and 5,422,251. Proteins include, but are not limited to, DNA-binding proteins, transcription factors, chromatin remodeling factors, methylated DNA binding proteins, polymerases, methylases, demethylases, acetylases, deacetylases, kinases, phosphatases, integrases, recombinases, ligases, topoisomerases, gyrases and helicases.

[0074] An exogenous molecule can be the same type of molecule as an endogenous molecule, e.g., an exogenous protein or nucleic acid. For example, an exogenous nucleic acid can comprise an infecting viral genome, a plasmid or episome introduced into a cell, or a chromosome that is not normally present in the cell. Methods for the introduction of exogenous molecules into cells are known to those of skill in the art and include, but are not limited to, lipid-mediated transfer (i.e., liposomes, including neutral and cationic lipids), electroporation, direct injection, cell fusion, particle bombardment, calcium phosphate co-precipitation, DEAE-dextran-mediated transfer and viral vector-mediated transfer. An exogenous molecule can also be the same type of molecule as an endogenous molecule but derived from a different species than the cell is derived from. For example, a human nucleic acid sequence may be introduced into a cell line originally derived from a mouse or hamster.

[0075] By contrast, an "endogenous" molecule is one that is normally present in a particular cell at a particular developmental stage under particular environmental conditions. For example, an endogenous nucleic acid can comprise a chromosome, the genome of a mitochondrion, chloroplast or other organelle, or a naturally-occurring episomal nucleic acid. Additional endogenous molecules can include proteins, for example, transcription factors and enzymes.

[0076] As used herein, the term "product of an exogenous nucleic acid" includes both polynucleotide and polypeptide products, for example, transcription products (polynucleotides such as RNA) and translation products (polypeptides).

[0077] A "fusion" molecule is a molecule in which two or more subunit molecules are linked, preferably covalently. The subunit molecules can be the same chemical type of molecule, or can be different chemical types of molecules. Examples of the first type of fusion molecule include, but are not limited to, fusion proteins (for example, a fusion between a ZFP or TALE DNA-binding domain and one or more activation domains) and fusion nucleic acids (for example, a nucleic acid encoding the fusion protein described *supra*). Examples of the second type of fusion molecule include, but are not limited to, a fusion between a triplex-forming nucleic acid and a polypeptide, and a fusion between a minor groove binder and a nucleic acid.

[0078] Expression of a fusion protein in a cell can result from delivery of the fusion protein to the cell or by delivery of a polynucleotide encoding the fusion protein to a cell, wherein the polynucleotide is transcribed, and the transcript is translated, to generate the fusion protein. Trans-splicing, polypeptide cleavage and polypeptide ligation can also be involved in expression of a protein in a cell. Methods for polynucleotide and polypeptide delivery to cells are presented elsewhere in this disclosure.

[0079] A "gene," for the purposes of the present disclosure, includes a DNA region encoding a gene product (see *infra*), as well as all DNA regions which regulate the production of the gene product, whether or not such regulatory sequences are adjacent to coding sequences, transcribed sequences, and combinations of coding and transcribed sequences. Accordingly, a gene includes, but is not necessarily limited to, promoter sequences, terminators, translational regulatory sequences such as ribosome binding sites and internal ribosome entry sites, enhancers, silencers, insulators, boundary elements, replication origins, matrix

attachment sites and locus control regions.

[0080] "Gene expression" refers to the conversion of the information, contained in a gene, into a gene product. A gene product can be the direct transcriptional product of a gene (e.g., mRNA, tRNA, rRNA, antisense RNA, microRNA, ribozyme, structural RNA or any other type of RNA) or a protein produced by translation of an mRNA. Gene products also include RNAs which are modified, by processes such as capping, polyadenylation, methylation, and editing, and proteins modified by, for example, methylation, acetylation, phosphorylation, ubiquitination, ADP-ribosylation, myristilation, and glycosylation.

[0081] "Modulation" of gene expression refers to a change in the activity of a gene. Modulation of expression can include, but is not limited to, gene activation and gene repression. Genome editing (e.g., cleavage, alteration, inactivation, random mutation) can be used to modulate expression. Gene inactivation refers to any reduction in gene expression as compared to a cell that does not include a ZFP, TALE or CRISPR/Cas system as described herein. Thus, gene inactivation may be partial or complete.

[0082] A "protected" mRNA is one in which the mRNA has been altered in some manner to increase the stability or translation of the mRNA. Examples of protections include the use of replacement of up to 25% of the cytosine and uridine residues with 2-thiouridine (s2U) and 5-methylcytosine (m5C). The resulting mRNA exhibits less immunogenicity and more stability as compared with its unmodified counterpart. (see Karikó et al. ((2012), Molecular Therapy, Vol. 16, No. 11, pages 1833 - 1844). Other changes include the addition of a so-called ARCA cap, which increases the translationability of the *in vitro* produced mRNA (see U.S. Patent No. 7,074,596).

[0083] A "region of interest" is any region of cellular chromatin, such as, for example, a gene or a non-coding sequence within or adjacent to a gene, in which it is desirable to bind an exogenous molecule. Binding can be for the purposes of targeted DNA cleavage, targeted recombination, and combinations of targeted DNA cleavage and targeted recombination. A region of interest can be present in a chromosome, an episome, an organellar genome (e.g., mitochondrial, chloroplast), or an infecting viral genome, for example. A region of interest can be within the coding region of a gene, within transcribed non-coding regions such as, for example, leader sequences, trailer sequences or introns, or within non-transcribed regions, either upstream or downstream of the coding region. A region of interest can be as small as a single nucleotide pair or up to 2,000 nucleotide pairs in length, or any integral value of nucleotide pairs.

[0084] "Eukaryotic" cells include, but are not limited to, fungal cells (such as yeast), plant cells, animal cells, mammalian cells and human cells (e.g., T-cells).

[0085] The terms "operative linkage" and "operatively linked" (or "operably linked") are used interchangeably with reference to a juxtaposition of two or more components (such as sequence elements), in which the components are arranged such that both components function normally and allow the possibility that at least one of the components can mediate a function that is exerted upon at least one of the other components. By way of illustration, a transcriptional regulatory sequence, such as a promoter, is operatively linked to a coding sequence if the transcriptional regulatory sequence controls the level of transcription of the coding sequence in response to the presence or absence of one or more transcriptional regulatory factors. A transcriptional regulatory sequence is generally operatively linked in *cis* with a coding sequence, but need not be directly adjacent to it. For example, an enhancer is a transcriptional regulatory sequence that is operatively linked to a coding sequence, even though they are not contiguous.

[0086] With respect to fusion polypeptides, the term "operatively linked" can refer to the fact that each of the components performs the same function in linkage to the other component as it would if it were not so linked. For example, with respect to a fusion polypeptide in which a ZFP, TALE or Cas DNA-binding domain is fused to an activation domain, the ZFP, TALE or Cas DNA-binding domain and the activation domain are in operative

linkage if, in the fusion polypeptide, the ZFP, TALE or Cas DNA-binding domain portion is able to bind its target site, its binding site, and combinations of its target site and binding site, while the activation domain is able to upregulate gene expression. When a fusion polypeptide in which a ZFP, TALE or Cas DNA-binding domain is fused to a cleavage domain, the ZFP, TALE or Cas DNA-binding domain and the cleavage domain are in operative linkage if, in the fusion polypeptide, the ZFP, TALE or Cas DNA-binding domain portion is able to bind its target site, its binding site, and combinations of its target site and its binding site, while the cleavage domain is able to cleave DNA in the vicinity of the target site.

[0087] A "functional fragment" of a protein, polypeptide or nucleic acid is a protein, polypeptide or nucleic acid whose sequence is not identical to the full-length protein, polypeptide or nucleic acid, yet retains the same function as the full-length protein, polypeptide or nucleic acid. A functional fragment can possess more, fewer, or the same number of residues as the corresponding native molecule, can contain one or more amino acid or nucleotide substitutions, and can be combinations possessing more, fewer, or the same number of residues as the corresponding native molecule and containing one or more amino acid or nucleotide substitutions. Methods for determining the function of a nucleic acid (e.g., coding function, ability to hybridize to another nucleic acid) are well-known in the art. Similarly, methods for determining protein function are well-known. For example, the DNA-binding function of a polypeptide can be determined, for example, by filter-binding, electrophoretic mobility-shift, or immunoprecipitation assays. DNA cleavage can be assayed by gel electrophoresis. See Ausubel *et al.*, *supra*. The ability of a protein to interact with another protein can be determined, for example, by co-immunoprecipitation, two-hybrid assays or complementation, both genetic and biochemical.

[0088] A "vector" is capable of transferring gene sequences to target cells. Typically, "vector construct," "expression vector," and "gene transfer vector," mean any nucleic acid construct capable of directing the expression of a gene of interest and which can transfer gene sequences to target cells. Thus, the term includes cloning, and expression vehicles, as well as integrating vectors.

[0089] The terms "subject" and "patient" are used interchangeably and refer to mammals such as human patients and non-human primates, as well as experimental animals such as pigs, cows, rabbits, dogs, cats, rats, mice, and other animals. Accordingly, the term "subject" or "patient" as used herein means any mammalian patient or subject to which the or stem cells of the invention can be administered. Subjects of the present invention include those that have been exposed to one or more chemical toxins, including, for example, a nerve toxin.

[0090] "Stemness" refers to the relative ability of any cell to act in a stem cell-like manner, *i.e.*, the degree of toti-, pluri-, multi- or oligo-potency and expanded or indefinite self-renewal that any particular stem cell may have.

Nucleases

[0091] Described herein are compositions, particularly nucleases, that are useful for *in vivo* cleavage of a donor molecule carrying a transgene and nucleases for cleavage of the genome of a cell such that the transgene is integrated into the genome in a targeted manner. One or more of the nucleases may be naturally occurring. One or more of the nucleases may be non-naturally occurring, *i.e.*, engineered in the DNA-binding domain, the cleavage domain, and a combination of the DNA-binding domain and cleavage domain. For example, the DNA-binding domain of a naturally-occurring nuclease may be altered to bind to a selected target site (e.g., a meganuclease that has been engineered to bind to site different than the cognate binding site). In other aspects of the disclosure, the nuclease comprises heterologous DNA-binding and cleavage domains (e.g., zinc finger nucleases; TAL-effector domain DNA binding proteins; meganuclease DNA-binding domains with heterologous cleavage domains).

A. DNA-binding domains

[0092] The DNA binding domain of one or more of the nucleases used for *in vivo* cleavage, targeted cleavage of the genome of a cell, and combinations of *in vivo* cleavage and targeted cleavage of the genome of a cell may comprise a zinc finger protein. A single zinc finger protein is made up of multiple zinc finger domains (e.g., 3, 4, 5, 6, or more zinc finger domains). Each zinc finger domain of is about 30 amino acids in length that it contains a beta turn (containing the two zinc coordinating residues and an alpha helix (containing the two invariant zinc coordinating residues), which are held in a particular conformation that allow binding to the protein to a target sequence. Canonical (C2H2) zinc finger domains having two cysteine (Cys) zinc coordinating residues in the beta turn and two histidine (His) zinc coordinating residues in the alpha helix or non-canonical (CH3) can be used. See, e.g., U.S. Patent No. 9,234,187. A 7-amino acid recognition helix is contained between the zinc coordinating residues of the beta turn and the zinc coordinating residues of the alpha helix. The recognition helix region is numbered -1 to +6 within the zinc finger domain and the amino acids outside this recognition region (and excluding the zinc coordinating residues are referred to as backbone residues).

[0093] Preferably, the zinc finger protein is non-naturally occurring in that the recognition helix is engineered to bind to a target site of choice. See, for example, Beerli et al. (2002) Nature Biotechnol. 20 :135-141 ; Pabo et al. (2001) Ann. Rev. Biochem.70 :313-340 ; Isalan et al. (2001) Nature Biotechnol. 19 :656-660 ; Segal et al. (2001) Curr. Opin. Biotechnol. 12 :632-637 ; Choo et al. (2000) Curr. Opin. Struct. Biol. 10:411-416; U.S. Patent Nos. 6,453,242; 6,534,261; 6,599,692; 6,503,717; 6,689,558; 7,030,215; 6,794,136; 7,067,317; 7,262,054; 7,070,934; 7,361,635; 7,253,273; and U.S. Patent Publication Nos. 2005/0064474; 2007/0218528; 2005/0267061.

[0094] An engineered zinc finger binding domain can have a novel binding specificity, compared to a naturally-occurring zinc finger protein. Engineering methods include, but are not limited to, rational design and various types of selection. Rational design includes, for example, using databases comprising triplet (or quadruplet) nucleotide sequences and individual zinc finger amino acid sequences, in which each triplet or quadruplet nucleotide sequence is associated with one or more amino acid sequences of zinc fingers which bind the particular triplet or quadruplet sequence. See, for example, co-owned U.S. Patent Nos. 6,453,242 and 6,534,261.

[0095] The ZFP DNA-binding domains may further comprise on or more modifications to the backbone of one or more the component zinc finger domains. The specificity of a ZFP for a target DNA sequence is dependent upon sequence specific contacts between the zinc finger domains and specific DNA bases, in particular, between the recognition helix region and the target site (typically each recognition helix binds to a target subsite of 3 nucleotides). In addition, the zinc finger domains also comprise amino acid residues that take part in non-specific interactions with the phosphates of the DNA backbone. Elrod-Erickson et al. ((1996) Structure 4:1171) demonstrated through co-crystallization of a zinc finger protein and its cognate DNA target that there are specific amino acids capable of interacting with the phosphates on the DNA backbone through formation of hydrogen bonds. Zinc finger proteins that employ the well-known Zif268 backbone typically have an arginine as the amino terminal residue of their second strand of β -sheet, which is also the second position carboxyl-terminal to the second invariant cysteine. This position can be referred to as (-5) within each zinc finger domain, as it is 5th residue preceding the start of the α -helix (and position -5 relative to the recognition helix numbered -1 to +6). The arginine at this position can interact with a phosphate on the DNA backbone via formation of a charged hydrogen bond with its side-chain guanidinium group. Zinc finger proteins in the Zif268 backbone also frequently have a lysine at a position that is 4 residues amino-terminal to the first invariant cysteine. This position can be referred to as (-14) within each finger, as it is 14th residue preceding the start of

the α -helix for zinc fingers with two residues between the zinc coordinating cysteine residues (and position -14 relative to the recognition helix region numbered -1 to +6). The lysine can interact with a phosphate on the DNA backbone via formation of a water-mediated charged hydrogen bond with its side-chain amino group. Since phosphate groups are found all along the DNA backbone, this type of interaction between the zinc finger and a DNA molecule is generally considered to be non-sequence specific (J. Miller, Massachusetts Institute of Technology Ph.D. Thesis, 2002).

[0096] Recent studies have hypothesized that non-specific phosphate contacting side chains in some nucleases may also account for some amount of non-specificity of those nucleases (Kleinstiver et al. (2016) *Nature* 529(7587):490-5; Guilinger et al. (2014) *Nat Meth*: 429-435). Researchers have proposed that these nucleases may possess 'excess DNA-binding energy', meaning that the nucleases may have a greater affinity for their DNA target than is required to substantially bind and cleave the target site. Thus, attempts were made to decrease the cationic charges in the TALE DNA binding domain (Guilinger, *ibid*) or the Cas9 DNA binding domain (Kleinstiver, *ibid*) to lower the DNA-binding energy of these nucleases, which resulted in increased cleavage specificity *in vitro*. However, additional studies (Sternberg et al. (2015) *Nature* 527(7576): 110-113) also suggest a role in proper folding and activation of the Cas9 nuclease domain for some of the cationic amino acids that were mutated in the Kleinstiver study of the Cas9 DNA binding domain. Thus, the exact role of these amino acids in Cas9 activity is not known.

[0097] Disclosed are mutations to amino acids within the ZFP DNA binding domain ('ZFP backbone') that can interact non-specifically with phosphates on the DNA backbone, but they do not comprise changes in the DNA recognition helices. Disclosed are mutations of cationic amino acid residues in the ZFP backbone that are not required for nucleotide target specificity. These mutations in the ZFP backbone may comprise: mutating a cationic amino acid residue to a neutral or anionic amino acid residue; or mutating a polar amino acid residue to a neutral or non-polar amino acid residue. Mutations made at position (-5), position (-9), position (-14), and combinations of mutations selected from mutations made at position (-5), position (-9) and position (-14) relative to the DNA binding helix are preferred. A zinc finger may comprise one or more mutations at (-5), (-9), (-14), and combinations of mutations selected from mutations at (-5), (-9), and (-14). One or more zinc finger(s) in a multi-finger zinc finger protein may comprise mutations in (-5), (-9), (-14) and combinations selected from (-5), (-9), and (-14). The amino acids at (-5), (-9), (-14) and combinations selected from (-5), (-9), and (-14) (e.g. an arginine (R) or lysine (K)) may be mutated to an alanine (A), leucine (L), Ser (S), Asp (D), Glu (E), Tyr (Y) and/or glutamine (Q).

[0098] In any of these fusion polypeptides described herein, the ZFP partners may further comprise mutations in the zinc finger DNA binding domain in the (-5), (-9), (-14) positions, and combinations of mutations selected from mutations at (-5), (-9), and (-14). The Arg (R) at position -5 may be changed to a Tyr (Y), Asp (D), Glu (E), Leu (L), Gln (Q), or Ala (A). The Arg (R) at position (-9) may be replaced with Ser (S), Asp (D), or Glu (E). The Arg (R) at position (-14) may be replaced with Ser (S) or Gln (Q). The fusion polypeptides can comprise mutations in the zinc finger DNA binding domain where the amino acids at the (-5), (-9), (-14) positions, and combinations of mutations selected from mutations at (-5), (-9), and (-14) are changed to any of the above listed amino acids in any combination.

[0099] Exemplary selection methods, including phage display and two-hybrid systems, are disclosed in US Patents 5,789,538; 5,925,523; 6,007,988; 6,013,453; 6,410,248; 6,140,466; 6,200,759; and 6,242,568; as well as WO 98/37186; WO 98/53057; WO 00/27878; WO 01/88197 and GB 2,338,237. In addition, enhancement of binding specificity for zinc finger binding domains has been described, for example, in co-owned WO 02/077227.

[0100] Selection of target sites; ZFPs and methods for design and construction of fusion proteins (and polynucleotides encoding same) are known to those of skill in the art and described in detail in U.S. Patent Nos. 6,140,081; 5,789,538; 6,453,242; 6,534,261; 5,925,523; 6,007,988; 6,013,453; 6,200,759; PCT

Publication Nos. WO 95/19431; WO 96/06166; WO 98/53057; WO 98/54311; WO 00/27878; WO 01/60970; WO 01/88197; WO 02/099084; WO 98/53058; WO 98/53059; WO 98/53060; WO 02/016536 and WO 03/016496.

[0101] Nearly any linker (spacer) may be used between one or more of the components of the DNA-binding domain (e.g., zinc fingers), between one or more DNA-binding domains, between the DNA-binding domain and the functional domain (e.g., nuclease), and between one or more DNA-binding domains and between the DNA-binding domain and the functional domain. Non-limiting examples of suitable linker sequences include U.S. Patent Nos. 8,772,453; 7,888,121; 6,479,626; 6,903,185; and 7,153,949; and U.S. Patent Publication Nos. 2009/0305419; 2015/0064789 and 2015/0132269. Thus, the proteins described herein may include any combination of suitable linkers between the individual DNA-binding components, between the DNA-binding domain and the functional domain, or between one or more DNA-binding domains and between the DNA-binding domain and the functional domain of the compositions described herein.

B. Cleavage Domains

[0102] Any suitable cleavage domain can be operatively linked to the DNA-binding domains as described herein to form a nuclease. The cleavage domain may be heterologous to the DNA-binding domain, for example a zinc finger DNA-binding domain and a cleavage domain from a nuclease. Heterologous cleavage domains can be obtained from any endonuclease or exonuclease. Exemplary endonucleases from which a cleavage domain can be derived include, but are not limited to, restriction endonucleases and homing endonucleases. See, for example, 2002-2003 Catalogue, New England Biolabs, Beverly, MA; and Belfort et al. (1997) *Nucleic Acids Res.* 25:3379-3388. Additional enzymes which cleave DNA are known (e.g., S1 Nuclease; mung bean nuclease; pancreatic DNase I; micrococcal nuclease; yeast HO endonuclease; see also Linn et al. (eds.) *Nucleases*, Cold Spring Harbor Laboratory Press, 1993). One or more of these enzymes (or functional fragments thereof) can be used as a source of cleavage domains and cleavage half-domains.

[0103] Similarly, a cleavage half-domain can be derived from any nuclease or portion thereof, as set forth above, that requires dimerization for cleavage activity. In general, two fusion proteins are required for cleavage if the fusion proteins comprise cleavage half-domains. Alternatively, a single protein comprising two cleavage half-domains can be used. The two cleavage half-domains can be derived from the same endonuclease (or functional fragments thereof), or each cleavage half-domain can be derived from a different endonuclease (or functional fragments thereof). In addition, the target sites for the two fusion proteins are preferably disposed, with respect to each other, such that binding of the two fusion proteins to their respective target sites places the cleavage half-domains in a spatial orientation to each other that allows the cleavage half-domains to form a functional cleavage domain, e.g., by dimerizing. Thus, the near edges of the target sites may be separated by 5-8 nucleotides or by 15-18 nucleotides. However, any integral number of nucleotides or nucleotide pairs can intervene between two target sites (e.g., from 2 to 50 nucleotide pairs or more). In general, the site of cleavage lies between the target sites.

[0104] Restriction endonucleases (restriction enzymes) are present in many species and are capable of sequence-specific binding to DNA (at a recognition site), and cleaving DNA at or near the site of binding. Certain restriction enzymes (e.g., Type IIS) cleave DNA at sites removed from the recognition site and have separable binding and cleavage domains. For example, the Type IIS enzyme *Fok I* catalyzes double-stranded cleavage of DNA, at 9 nucleotides from its recognition site on one strand and 13 nucleotides from its recognition site on the other. See, for example, U.S. Patent Nos. 5,356,802; 5,436,150 and 5,487,994; as well as Li et al. (1992) *Proc. Natl. Acad. Sci. USA* 89:4275-4279; Li et al. (1993) *Proc. Natl. Acad. Sci. USA* 90:2764-2768; Kim et al. (1994a) *Proc. Natl. Acad. Sci. USA* 91:883-887; Kim et al. (1994b) *J. Biol. Chem.* 269:31,978-31,982. Thus, fusion proteins may comprise the cleavage domain (or cleavage half-domain) from at least one Type IIS restriction enzyme and one or more zinc finger binding domains, which may or may not

be engineered.

[0105] For optimal cleavage specificity by a sequence-selective (artificial) nuclease, it is desirable to arrange conditions so that on-target binding and activity is not saturating. Under saturating conditions - by definition - an excess of nuclease is used over what is necessary to achieve complete on-target activity. This excess provides no on-target benefit but can nonetheless result in increased cleavage at off-target sites. For monomeric nucleases, saturating conditions may be readily avoided by performing a simple dose response study to identify and avoid the saturating plateau on a titration curve. However, for a dimeric nuclease such as ZFN, TALEN or dCas-Fok, identifying and avoiding saturating conditions may be more complicated if the binding affinities of the individual monomers are dissimilar. In such cases, a dose response study using a simple 1:1 nuclease ratio will only reveal the saturation point of the weaker binding monomer. Under such a scenario, if, for example, monomer affinities differ by a factor of 10, then at the saturation point identified in a 1:1 titration study the higher affinity monomer will be present at a concentration that is 10-fold higher than it needs to be. The resulting excess of the higher affinity monomer can in turn lead to increased off-target activity without providing any beneficial increase in cleavage at the intended target, potentially leading to a decreased specificity overall for any given nuclease pair.

[0106] An exemplary Type IIS restriction enzyme, whose cleavage domain is separable from the binding domain, is *FokI*. This particular enzyme is active as a dimer. Bitinaite et al. (1998) Proc. Natl. Acad. Sci. USA 95:10,570-10,575. Accordingly, for the purposes of the present disclosure, the portion of the *FokI* enzyme used in the disclosed fusion proteins is considered a cleavage half-domain. Thus, for targeted double-stranded cleavage, targeted replacement of cellular sequences using zinc finger-*FokI* fusions, and combinations of targeted double-stranded cleavage and targeted replacement of cellular sequences using zinc finger-*FokI* fusions, two fusion proteins, each comprising a *FokI* cleavage half-domain, can be used to reconstitute a catalytically active cleavage domain. Alternatively, a single polypeptide molecule containing a zinc finger binding domain and two *FokI* cleavage half-domains can also be used. Parameters for targeted cleavage and targeted sequence alteration using zinc finger-*FokI* fusions are provided elsewhere in this disclosure.

[0107] A cleavage domain or cleavage half-domain can be any portion of a protein that retains cleavage activity, or that retains the ability to multimerize (e.g., dimerize) to form a functional cleavage domain.

[0108] Exemplary Type IIS restriction enzymes are described in U.S. Patent No. 7,888,121. Additional restriction enzymes also contain separable binding and cleavage domains, and these are contemplated by the present disclosure. See, for example, Roberts et al. (2003) Nucleic Acids Res. 31:418-420.

[0109] The cleavage domain may comprise one or more engineered cleavage half-domain (also referred to as dimerization domain mutants) that minimize or prevent homodimerization, as described, for example, in See, e.g., U.S. Patent Nos. 7,914,796; 8,034,598 and 8,623,618. Amino acid residues at positions 446, 447, 479, 483, 484, 486, 487, 490, 491, 496, 498, 499, 500, 531, 534, 537, and 538 of *FokI* are all targets for influencing dimerization of the *FokI* cleavage half-domains where the numbering is with respect to the crystal structures 1FOK.pdb and 2FOK.pdb (see Wah et al. (1997) Nature 388:97-100) having the sequence shown below:

Wild	type	FokI	cleavage	half	domain	(SEQ	ID	NO:18)
QLVKSELEEKKSEL	RHKLKYP	HEYIELIEIARN	STQDRILEMKV	MEFFMKVY				
GYRGKHLGGSRK	PDGAIYTVG	SPIDYGVIVDT	KAYSGGYNLP	IGQADEMQR				
VEENQTRNKHIN	PNEWWKVYP	SSVTEFKFLF	VSGHFKGNY	KAQLTRLN				
CNGAVLSVEELL	GEMIKAGT	LTLEEVRRK	FNNGEINF					

[0110] Exemplary engineered cleavage half-domains of *FokI* that form obligate heterodimers include a pair in which a first cleavage half-domain includes mutations at amino acid residues at positions 490 and 538 of *FokI* and a second cleavage half-domain includes mutations at amino acid residues 486 and 499.

[0111] Thus, a mutation at 490 may replace Glu (E) with Lys (K); the mutation at 538 may replace Ile (I) with Lys (K); the mutation at 486 may replace Gln (Q) with Glu (E); and the mutation at position 499 may replace Ile (I) with Lys (K). Specifically, the engineered cleavage half-domains described herein were prepared by mutating positions 490 (E→K) and 538 (I→K) in one cleavage half-domain to produce an engineered cleavage half-domain designated "E490K:I538K" and by mutating positions 486 (Q→E) and 499 (I→L) in another cleavage half-domain to produce an engineered cleavage half-domain designated "Q486E:I499L". The engineered cleavage half-domains described herein are obligate heterodimer mutants in which aberrant cleavage via the ZFN homodimers is minimized or abolished. See, e.g., U.S. Patent Publication No. 2008/0131962. The engineered cleavage half-domain may comprise mutations at positions 486, 499 and 496 (numbered relative to wild-type FokI), for instance mutations that replace the wild type Gln (Q) residue at position 486 with a Glu (E) residue, the wild type Ile (I) residue at position 499 with a Leu (L) residue and the wild-type Asn (N) residue at position 496 with an Asp (D) or Glu (E) residue (also referred to as a "ELD" and "ELE" domains, respectively). The engineered cleavage half-domain may comprise mutations at positions 490, 538 and 537 (numbered relative to wild-type FokI), for instance mutations that replace the wild type Glu (E) residue at position 490 with a Lys (K) residue, the wild type Ile (I) residue at position 538 with a Lys (K) residue, and the wild-type His (H) residue at position 537 with a Lys (K) residue or a Arg (R) residue (also referred to as "KKK" and "KKR" domains, respectively). The engineered cleavage half-domain may comprise mutations at positions 490 and 537 (numbered relative to wild-type FokI), for instance mutations that replace the wild type Glu (E) residue at position 490 with a Lys (K) residue and the wild-type His (H) residue at position 537 with a Lys (K) residue or a Arg (R) residue (also referred to as "KIK" and "KIR" domains, respectively). See, e.g., U.S. Patent Nos. 7,914,796; 8,034,598 and 8,623,618. The engineered cleavage half domain may comprise the "Sharkey", "Sharkey" mutations, and combinations of the "Sharkey" and "Sharkey" mutations (see Guo et al. (2010) J. Mol. Biol. 400(1):96-107).

[0112] Thus, cleavage half domains derived from FokI may comprise a mutation in one or more of amino acid residues as shown in SEQ ID NO:18, including mutations in the dimerization domain as described above; mutations in the catalytic domain, mutations in other amino acid residues such as phosphate contact residues, and any combination of mutations selected from mutations in the dimerization domain, mutations in the catalytic domain, and mutations in other amino acid residues such as phosphate contact residues. Mutations include substitutions (of a wild-type amino acid residue with a different residue), insertions (of one or more amino acid residues), deletions (of one or more amino acid residues), and any combination of mutations selected from substitutions, insertions and deletions. One or more of residues 414-426, 443-450, 467-488, 501-502, 521-531 (numbered relative to SEQ ID NO:18), and any combination of such residues, may be mutated since these residues are located close to the DNA backbone in a molecular model of a ZFN bound to its target site described in Miller et al. ((2007) Nat Biotechnol 25:778-784). One or more residues at positions 416, 422, 447, 448, and 525 may be mutated. The mutation may comprise a substitution of a wild-type residue with a different residue, for example a serine (S) residue. The FokI cleavage domain of the nucleases described herein may comprise an ELD dimerization domain mutation, a KKR dimerization domain mutation, a K525S mutation, and any combination selected from an ELD dimerization domain mutation or a KKR dimerization domain mutation and a K525S mutation.

[0113] Engineered cleavage domains described herein can be prepared using any suitable method, for example, by site-directed mutagenesis of wild-type cleavage half-domains (*Fok I*) as described in U.S. Patent Nos. 7,888,121; 7,914,796; 8,034,598 and 8,623,618. Furthermore, the cleavage domains described herein may be fused to a DNA-binding domain (e.g., ZFP) using any suitable linker, including, but not limited to the linkers described in U.S. Patent Nos. 9,394,531 and 9,567,609.

[0114] Alternatively, nucleases may be assembled *in vivo* at the nucleic acid target site using so-called "split-enzyme" technology (see, e.g. U.S. Patent Publication No. 2009/0068164). Components of such split enzymes may be expressed either on separate expression constructs, or can be linked in one open reading frame where the individual components are separated, for example, by a self-cleaving 2A peptide or IRES sequence.

Components may be individual zinc finger binding domains or domains of a meganuclease nucleic acid binding domain.

[0115] Nucleases can be screened for activity prior to use, for example in a yeast-based chromosomal system as described in WO 2009/042163 and 20090068164. Expression of the nuclease may be under the control of a constitutive promoter or an inducible promoter, for example the galactokinase promoter which is activated (de-repressed) in the presence of raffinose, galactose and a combination of raffinose and galactose and repressed in presence of glucose.

[0116] The nuclease(s) as described herein may make one or more double-stranded, one or more single-stranded cuts, and combinations of one or more double-stranded and one or more single-stranded cuts in the target site. In certain aspects of the disclosure, the nuclease comprises a catalytically inactive cleavage domain (e.g., FokI, a Cas protein and combinations of FokI and a Cas protein). See, e.g., U.S. Patent Nos. 9,200,266; 8,703,489 and Guillingier et al. (2014) Nature Biotech. 32(6):577-582. The catalytically inactive cleavage domain may, in combination with a catalytically active domain act as a nickase to make a single-stranded cut. Therefore, two nickases can be used in combination to make a double-stranded cut in a specific region. Additional nickases are also known in the art, for example, McCaffery et al. (2016) Nucleic Acids Res. 44(2):e11. doi: 10.1093/nar/gkv878. Epub 2015 Oct 19.

Target Sites

[0117] As described in detail above, DNA domains can be engineered to bind to any sequence of choice. An engineered DNA-binding domain can have a novel binding specificity, compared to a naturally-occurring DNA-binding domain. The DNA-binding domains may bind to a sequence within a BCL11A enhancer sequence, for example a target site (typically 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21 or even more base pairs) is between exon 2 and exon 3 of BCL11A, including DNA-binding domains that bind to a sequence within a DNaseI hypersensitive site in the BCL11A enhancer sequence (e.g., +58) as shown in Table 1. Engineering methods include, but are not limited to, rational design and various types of selection. Rational design includes, for example, using databases comprising triplet (or quadruplet) nucleotide sequences and individual zinc finger amino acid sequences, in which each triplet or quadruplet nucleotide sequence is associated with one or more amino acid sequences of zinc fingers which bind the particular triplet or quadruplet sequence. See, for example, co-owned U.S. Patent Nos. 6,453,242 and 6,534,261. Rational design of TAL-effector domains can also be performed. See, e.g., U.S. Patent Publication No. 2011/0301073.

[0118] Exemplary selection methods applicable to DNA-binding domains, including phage display and two-hybrid systems, are disclosed in U.S. Patent Nos. 5,789,538; 5,925,523; 6,007,988; 6,013,453; 6,410,248; 6,140,466; 6,200,759; and 6,242,568; as well as PCT Publication Nos. WO 98/37186; WO 98/53057; WO 00/27878; WO 01/88197 and U.K. Patent No. GB 2,338,237. In addition, enhancement of binding specificity for zinc finger binding domains has been described, for example, in co-owned WO 02/077227.

[0119] Selection of target sites; nucleases and methods for design and construction of fusion proteins (and polynucleotides encoding same) are known to those of skill in the art and described in detail in U.S. Patent Publication Nos. 2005/0064474 and 2006/0188987.

[0120] In addition, as disclosed in these and other references, DNA-binding domains (e.g., multi-fingered zinc finger proteins) and fusions of DNA-binding domain(s) and functional domain(s) may be linked together using any suitable linker sequences, including for example, linkers of 5 or more amino acids. U.S. Patent Nos. 8,772,453; 7,888,121 (e.g., "ZC" linker); 6,479,626; 6,903,185; and 7,153,949; U.S. Publication No. 2009/0305419) and 2015/0064789. The proteins described herein may include any combination of suitable linkers between the individual DNA-binding domains of the protein. See, also, U.S. Patent No. 8,586,526.

Donors

[0121] The present disclosure relates to nuclease-mediated targeted integration of an exogenous sequence into the genome of a cell using the BCL11A enhancer region-binding molecules described herein. As noted above, insertion of an exogenous sequence (also called a "donor sequence" or "donor" or "transgene"), for example for deletion of a specified region, for correction of a mutant gene, for a combination of deletion of a specified region and correction of a mutant gene, or for increased expression of a wild-type gene. It will be readily apparent that the donor sequence is typically not identical to the genomic sequence where it is placed. A donor sequence can contain a non-homologous sequence flanked by two regions of homology to allow for efficient HDR at the location of interest or can be integrated via non-homology directed repair mechanisms such as NHEJ. Additionally, donor sequences can comprise a vector molecule containing sequences that are not homologous to the region of interest in cellular chromatin. A donor molecule can contain several, discontinuous regions of homology to cellular chromatin, and, for example, lead to a deletion of a BCL11A enhancer region (or a fragment thereof) when used as a substrate for repair of a DSB induced by one of the nucleases described here. Further, for targeted insertion of sequences not normally present in a region of interest, said sequences can be present in a donor nucleic acid molecule and flanked by regions of homology to sequence in the region of interest.

[0122] Polynucleotides for insertion can also be referred to as "exogenous" polynucleotides, "donor" polynucleotides or molecules or "transgenes." The donor polynucleotide can be DNA or RNA, single-stranded or double-stranded, and can be introduced into a cell in linear or circular form. See, e.g., U.S. Patent Application Publication Nos. 2010/0047805 and 2011/0207221. The donor sequence(s) are preferably contained within a DNA MC, which may be introduced into the cell in circular or linear form. If introduced in linear form, the ends of the donor sequence can be protected (e.g., from exonucleolytic degradation) by methods known to those of skill in the art. For example, one or more dideoxynucleotide residues are added to the 3' terminus of a linear molecule and self-complementary oligonucleotides are optionally ligated to one or both ends. See, for example, Chang et al. (1987) Proc. Natl. Acad. Sci. USA 84:4959-4963; Nehls et al. (1996) Science 272:886-889. Additional methods for protecting exogenous polynucleotides from degradation include, but are not limited to, addition of terminal amino group(s) and the use of modified internucleotide linkages such as, for example, phosphorothioates, phosphoramidates, and O-methyl ribose or deoxyribose residues. If introduced in double-stranded form, the donor may include one or more nuclease target sites, for example, nuclease target sites flanking the transgene to be integrated into the cell's genome. See, e.g., U.S. Patent Publication No. 2013/0326645.

[0123] A polynucleotide can be introduced into a cell as part of a vector molecule having additional sequences such as, for example, replication origins, promoters and genes encoding antibiotic resistance. Moreover, donor polynucleotides can be introduced as naked nucleic acid, as nucleic acid complexed with an agent such as a liposome or poloxamer, or can be delivered by viruses (e.g., adenovirus, AAV, herpesvirus, retrovirus, lentivirus and integrase defective lentivirus (IDLV)).

[0124] In certain embodiments, the double-stranded donor includes sequences (e.g., coding sequences, also referred to as transgenes) greater than 1 kb in length, for example between 2 and 200 kb, between 2 and 10 kb (or any value therebetween). The double-stranded donor also includes at least one nuclease target site, for example. In certain embodiments, the donor includes at least 2 target sites, for example for a pair of ZFNs or TALENs. Typically, the nuclease target sites are outside the transgene sequences, for example, 5' and/or 3' to the transgene sequences, for cleavage of the transgene. The nuclease cleavage site(s) may be for any nuclease(s). In certain embodiments, the nuclease target site(s) contained in the double-stranded donor are for the same nuclease(s) used to cleave the endogenous target into which the cleaved donor is integrated via homology-independent methods.

[0125] The donor is generally inserted so that its expression is driven by the endogenous promoter at the integration site, namely the promoter that drives expression of the endogenous gene into which the donor is inserted (e.g., globin, AAVS1, etc.). However, it will be apparent that the donor may comprise a promoter, an enhancer, and combinations of both a promoter and enhancer, for example a constitutive promoter or an inducible or tissue specific promoter.

[0126] The donor molecule may be inserted into an endogenous gene such that all, some or none of the endogenous gene is expressed. In other embodiments, the transgene (e.g., with or without globin encoding sequences) is integrated into any endogenous locus, for example a safe-harbor locus. See, e.g., U.S. Patent Publication Nos. 2008/0299580; 2008/0159996 and 2010/00218264.

[0127] Furthermore, although not required for expression, exogenous sequences may also include transcriptional or translational regulatory sequences, for example, promoters, enhancers, insulators, internal ribosome entry sites, sequences encoding 2A peptides, polyadenylation signals, and combinations thereof.

[0128] The transgenes carried on the donor sequences described herein may be isolated from plasmids, cells or other sources using standard techniques known in the art such as PCR. Donors for use can include varying types of topology, including circular supercoiled, circular relaxed, linear and the like. Alternatively, they may be chemically synthesized using standard oligonucleotide synthesis techniques. In addition, donors may be methylated or lack methylation. Donors may be in the form of bacterial or yeast artificial chromosomes (BACs or YACs).

[0129] The double-stranded donor polynucleotides described herein may include one or more non-natural bases, one or more backbones, and combinations of one or more non-natural bases and one or more backbones. In particular, insertion of a donor molecule with methylated cytosines may be carried out using the methods described herein to achieve a state of transcriptional quiescence in a region of interest.

[0130] The exogenous (donor) polynucleotide may comprise any sequence of interest (exogenous sequence). Exemplary exogenous sequences include, but are not limited to any polypeptide coding sequence (e.g., cDNAs), promoter sequences, enhancer sequences, epitope tags, marker genes, cleavage enzyme recognition sites and various types of expression constructs. Marker genes include, but are not limited to, sequences encoding proteins that mediate antibiotic resistance (e.g., ampicillin resistance, kanamycin resistance, neomycin resistance, G418 resistance, puromycin resistance, hygromycin resistance, blasticidin resistance), sequences encoding colored or fluorescent or luminescent proteins (e.g., green fluorescent protein, enhanced green fluorescent protein, red fluorescent protein, luciferase), and proteins which mediate enhanced cell growth, gene amplification (e.g., dihydrofolate reductase) and combinations of enhanced cell growth and gene amplification. Epitope tags include, for example, one or more copies of FLAG, His, Myc, tandem affinity purification (TAP), HA, biotinylatable peptide, or any detectable amino acid sequence.

[0131] In a preferred embodiment, the exogenous sequence (transgene) comprises a polynucleotide encoding any polypeptide of which expression in the cell is desired, including, but not limited to antibodies, antigens, enzymes, receptors (cell surface or nuclear), hormones, lymphokines, cytokines, reporter polypeptides, growth factors, and functional fragments of any of the above. The coding sequences may be, for example, cDNAs.

[0132] For example, the exogenous sequence may comprise a sequence encoding a polypeptide that is lacking or non-functional in the subject having a genetic disease, including but not limited to any of the following genetic diseases: achondroplasia, achromatopsia, acid maltase deficiency, adenosine deaminase deficiency (OMIM No. 102700), adrenoleukodystrophy, aicardi syndrome, alpha-1 antitrypsin deficiency, alpha-thalassemia, androgen insensitivity syndrome, apert syndrome, arrhythmogenic right ventricular, dysplasia,

ataxia telangiectasia, Barth syndrome, beta-thalassemia, blue rubber bleb nevus syndrome, Canavan disease, chronic granulomatous diseases (CGD), cri du chat syndrome, cystic fibrosis, Dercum's disease, ectodermal dysplasia, Fanconi anemia, fibrodysplasia ossificans progressiva, fragile X syndrome, galactosemia, Gaucher's disease, generalized gangliosidosis (e.g., GM1), hemochromatosis, the hemoglobin C mutation in the 6th codon of beta-globin (HbC), hemophilia, Huntington's disease, Hurler Syndrome, hypophosphatasia, Klinefelter syndrome, Krabbe Disease, Langer-Giedion Syndrome, leukocyte adhesion deficiency (LAD, OMIM No. 116920), leukodystrophy, long QT syndrome, Marfan syndrome, Moebius syndrome, mucopolysaccharidosis (MPS), nail patella syndrome, nephrogenic diabetes insipidus, neurofibromatosis, Niemann-Pick disease, osteogenesis imperfecta, porphyria, Prader-Willi syndrome, progeria, Proteus syndrome, retinoblastoma, Rett syndrome, Rubinstein-Taybi syndrome, Sanfilippo syndrome, severe combined immunodeficiency (SCID), Shwachman syndrome, sickle cell disease (sickle cell anemia), Smith-Magenis syndrome, Stickler syndrome, Tay-Sachs disease, Thrombocytopenia Absent Radius (TAR) syndrome, Treacher Collins syndrome, trisomy, tuberous sclerosis, Turner's syndrome, urea cycle disorder, von Hippel-Landau disease, Waardenburg syndrome, Williams syndrome, Wilson's disease, Wiskott-Aldrich syndrome, X-linked lymphoproliferative syndrome (XLP, OMIM No. 308240).

[0133] Additional exemplary diseases that can be treated by targeted integration include acquired immunodeficiencies, lysosomal storage diseases (e.g., Gaucher's disease, GM1, Fabry disease and Tay-Sachs disease), mucopolysaccharidosis (e.g. Hunter's disease, Hurler's disease), hemoglobinopathies (e.g., sickle cell disease, HbC, α -thalassemia, β -thalassemia) and hemophilias.

[0134] In certain embodiments, the exogenous sequences can comprise a marker gene (described above), allowing selection of cells that have undergone targeted integration, and a linked sequence encoding an additional functionality. Non-limiting examples of marker genes include GFP, drug selection marker(s) and the like.

[0135] Additional gene sequences that can be inserted may include, for example, wild-type genes to replace mutated sequences. For example, a wild-type Factor IX gene sequence may be inserted into the genome of a stem cell in which the endogenous copy of the gene is mutated. The wild-type copy may be inserted at the endogenous locus, or may alternatively be targeted to a safe harbor locus.

[0136] Construction of such expression cassettes, following the teachings of the present specification, utilizes methodologies well known in the art of molecular biology (see, for example, Ausubel or Maniatis). Before use of the expression cassette to generate a transgenic animal, the responsiveness of the expression cassette to the stress-inducer associated with selected control elements can be tested by introducing the expression cassette into a suitable cell line (e.g., primary cells, transformed cells, or immortalized cell lines).

[0137] Furthermore, although not required for expression, exogenous sequences may also transcriptional or translational regulatory sequences, for example, promoters, enhancers, insulators, internal ribosome entry sites, sequences encoding 2A peptides, polyadenylation signals, and combinations of 2A polypeptides and polyadenylation signals. Further, the control elements of the genes of interest can be operably linked to reporter genes to create chimeric genes (e.g., reporter expression cassettes).

[0138] Targeted insertion of non-coding nucleic acid sequence may also be achieved. Sequences encoding antisense RNAs, RNAi, shRNAs and micro RNAs (miRNAs) may also be used for targeted insertions.

[0139] In additional embodiments, the donor nucleic acid may comprise non-coding sequences that are specific target sites for additional nuclease designs. Subsequently, additional nucleases may be expressed in cells such that the original donor molecule is cleaved and modified by insertion of another donor molecule of interest. In this way, reiterative integrations of donor molecules may be generated allowing for trait stacking at a particular locus of interest or at a safe harbor locus.

Delivery

[0140] The nucleases as described herein (Table 1), polynucleotides encoding these nucleases, donor polynucleotides and compositions comprising the proteins, polynucleotides, and combinations of proteins and polynucleotides described herein may be delivered *in vivo* or *ex vivo* by any suitable means into any cell type.

[0141] Suitable cells include eukaryotic (e.g., animal) and prokaryotic cells and eukaryotic and prokaryotic cell lines. Non-limiting examples of such cells or cell lines generated from such cells include COS, CHO (e.g., CHO-S, CHO-K1, CHO-DG44, CHO-DUXB11, CHO-DUKX, CHOK1SV), VERO, MDCK, WI38, V79, B14AF28-G3, BHK, HaK, NS0, SP2/0-Ag14, HeLa, HEK293 (e.g., HEK293-F, HEK293-H, HEK293-T), and perC6 cells as well as insect cells such as *Spodoptera frugiperda* (Sf), or fungal cells such as *Saccharomyces*, *Pichia* and *Schizosaccharomyces*. In certain embodiments, the cell line is a CHO, MDCK or HEK293 cell line. Suitable cells also include stem cells such as, by way of example, embryonic stem cells, induced pluripotent stem cells, hematopoietic stem cells, neuronal stem cells and mesenchymal stem cells.

[0142] Methods of delivering nucleases as described herein are described, for example, in U.S. Patent Nos. 6,453,242; 6,503,717; 6,534,261; 6,599,692; 6,607,882; 6,689,558; 6,824,978; 6,933,113; 6,979,539; 7,013,219; and 7,163,824.

[0143] Nucleases, donor constructs, and combinations of nucleases and donor constructs as described herein may also be delivered using vectors containing sequences encoding one or more of the ZFN(s), described herein. Any vector systems may be used including, but not limited to, plasmid vectors, retroviral vectors, lentiviral vectors, adenovirus vectors, poxvirus vectors; herpesvirus vectors and adeno-associated virus vectors, etc. See, also, U.S. Patent Nos. 6,534,261; 6,607,882; 6,824,978; 6,933,113; 6,979,539; 7,013,219; and 7,163,824. Furthermore, it will be apparent that any of these vectors may comprise one or more of the sequences needed for treatment. Thus, when one or more nucleases and a donor construct are introduced into the cell, the nucleases, donor polynucleotide, and combinations of nucleases and donor polynucleotide may be carried on the same vector or on different vectors (DNA MC(s)). When multiple vectors are used, each vector may comprise a sequence encoding one or multiple nucleases, one or more donor constructs, and combinations of one or more nucleases and one or more donor constructions. Conventional viral and non-viral based gene transfer methods can be used to introduce nucleic acids encoding nucleases, donor constructs, and combinations of nucleases and donor constructs in cells (e.g., mammalian cells) and target tissues. Non-viral vector delivery systems include DNA or RNA plasmids, DNA MCs, naked nucleic acid, and nucleic acid complexed with a delivery vehicle such as a liposome or poloxamer. Suitable non-viral vectors include nanotaxis vectors, including vectors commercially available from InCellArt (France). Viral vector delivery systems include DNA and RNA viruses, which have either episomal or integrated genomes after delivery to the cell. For a review of *in vivo* delivery of engineered DNA-binding proteins and fusion proteins comprising these binding proteins, see, e.g., Rebar (2004) Expert Opinion Invest. Drugs 13(7):829-839; Rossi et al. (2007) Nature Biotech. 25(12): 1444-1454 as well as general gene delivery references such as Anderson, Science 256:808-813 (1992); Nabel & Felgner, TIBTECH 11:211-217 (1993); Mitani & Caskey, TIBTECH 11:162-166 (1993); Dillon, TIBTECH 11:167-175 (1993); Miller, Nature 357:455-460 (1992); Van Brunt, Biotechnology 6(10): 1149-1154 (1988); Vigne, Restorative Neurology and Neuroscience 8:35-36 (1995); Kremer & Perricaudet, British Medical Bulletin 51(1):31-44 (1995); Haddada et al., in Current Topics in Microbiology and Immunology Doerfler and Böhm (eds.) (1995); and Yu et al., Gene Therapy 1:13-26 (1994).

[0144] Methods of non-viral delivery of nucleic acids include electroporation, lipofection, microinjection, biolistics, virosomes, liposomes, immunoliposomes, polycation or lipid:nucleic acid conjugates, naked DNA, artificial virions, membrane deformation, and agent-enhanced uptake of DNA. Sonoporation using, e.g., the Sonitron 2000 system (Rich-Mar) can also be used for delivery of nucleic acids.

[0145] Additional exemplary nucleic acid delivery systems include those provided by Amaxa Biosystems (Cologne, Germany), Maxcyte, Inc. (Rockville, Maryland), BTX Molecular Delivery Systems (Holliston, MA) and Copernicus Therapeutics Inc., (see for example US6008336). Lipofection is described in e.g., U.S. Patent Nos. 5,049,386; 4,946,787; and 4,897,355) and lipofection reagents are sold commercially (e.g., Transfectam™ and Lipofectin™). Cationic and neutral lipids that are suitable for efficient receptor-recognition lipofection of polynucleotides include those of Felgner, WO 91/17424, WO 91/16024.

[0146] The preparation of lipid:nucleic acid complexes, including targeted liposomes such as immunolipid complexes, is well known to one of skill in the art (see, e.g., Crystal, Science 270:404-410 (1995); Blaese et al., Cancer Gene Ther. 2:291-297 (1995); Behr et al., Bioconjugate Chem. 5:382-389 (1994); Remy et al., Bioconjugate Chem. 5:647-654 (1994); Gao et al., Gene Therapy 2:710-722 (1995); Ahmad et al., Cancer Res. 52:4817-4820 (1992); U.S. Pat. Nos. 4,186,183, 4,217,344, 4,235,871, 4,261,975, 4,485,054, 4,501,728, 4,774,085, 4,837,028, and 4,946,787). Other lipid:nucleic acid complexes include those comprising novel cationic lipids, novel pegylated lipids, and combinations of novel cationic lipids and novel pegylated lipids (see e.g. U.S. Provisional Patent Application Nos. 62/432,042 and 62/458,373).

[0147] Additional methods of delivery include the use of packaging the nucleic acids to be delivered into EnGeneIC delivery vehicles (EDVs). These EDVs are specifically delivered to target tissues using bispecific antibodies where one arm of the antibody has specificity for the target tissue and the other has specificity for the EDV. The antibody brings the EDVs to the target cell surface and then the EDV is brought into the cell by endocytosis. Once in the cell, the contents are released (see MacDiarmid et al. (2009) Nature Biotechnology 27(7):643).

[0148] The use of RNA or DNA viral based systems for the delivery of nucleic acids encoding engineered ZFPs, TALEs and CRISPR/Cas systems take advantage of highly evolved processes for targeting a virus to specific cells in the body and trafficking the viral payload to the nucleus. Viral vectors can be administered directly to patients (*in vivo*) or they can be used to treat cells *in vitro* and the modified cells are administered to patients (*ex vivo*). Conventional viral based systems for the delivery of ZFPs include, but are not limited to, retroviral, lentiviral, adenoviral, adeno-associated, vaccinia and herpes simplex virus vectors for gene transfer. Integration in the host genome is possible with the retrovirus, lentivirus, and adeno-associated virus gene transfer methods, often resulting in long term expression of the inserted transgene. Additionally, high transduction efficiencies have been observed in many different cell types and target tissues.

[0149] The tropism of a retrovirus can be altered by incorporating foreign envelope proteins, expanding the potential target population of target cells. Lentiviral vectors are retroviral vectors that are able to transduce or infect non-dividing cells and typically produce high viral titers. Selection of a retroviral gene transfer system depends on the target tissue. Retroviral vectors are comprised of *cis*-acting long terminal repeats with packaging capacity for up to 6-10 kb of foreign sequence. The minimum *cis*-acting LTRs are sufficient for replication and packaging of the vectors, which are then used to integrate the therapeutic gene into the target cell to provide permanent transgene expression. Widely used retroviral vectors include those based upon murine leukemia virus (MuLV), gibbon ape leukemia virus (GaLV), Simian Immunodeficiency virus (SIV), human immunodeficiency virus (HIV), and combinations thereof (see, e.g., Buchscher et al., J. Virol. 66:2731-2739 (1992); Johann et al., J. Virol. 66:1635-1640 (1992); Sommerfelt et al., Virol. 176:58-59 (1990); Wilson et al., J. Virol.63:2374-2378 (1989); Miller et al., J. Virol. 65:2220-2224 (1991); PCT/US94/05700).

[0150] In applications in which transient expression is preferred, adenoviral based systems can be used. Adenoviral based vectors are capable of very high transduction efficiency in many cell types and do not require cell division. With such vectors, high titer and high levels of expression have been obtained. This vector can be produced in large quantities in a relatively simple system. Adeno-associated virus ("AAV") vectors are also used to transduce cells with target nucleic acids, e.g., in the *in vitro* production of nucleic

acids and peptides, and for *in vivo* and *ex vivo* gene therapy procedures (see, e.g., West et al., Virology 160:38-47 (1987); U.S. Patent No. 4,797,368; WO 93/24641; Kotin, Human Gene Therapy 5:793-801 (1994); Muzyczka, J. Clin. Invest. 94:1351 (1994). Construction of recombinant AAV vectors are described in a number of publications, including U.S. Pat. No. 5,173,414; Tratschin et al., Mol. Cell. Biol. 5:3251-3260 (1985); Tratschin, et al., Mol. Cell. Biol. 4:2072-2081 (1984); Hermonat & Muzyczka, PNAS 81:6466-6470 (1984); and Samulski et al., J. Virol. 63:03822-3828 (1989).

[0151] At least six viral vector approaches are currently available for gene transfer in clinical trials, which utilize approaches that involve complementation of defective vectors by genes inserted into helper cell lines to generate the transducing agent.

[0152] pLASN and MFG-S are examples of retroviral vectors that have been used in clinical trials (Dunbar et al., Blood 85:3048-305 (1995); Kohn et al., Nat. Med. 1:1017-102 (1995); Malech et al., PNAS 94:22 12133-12138 (1997)). PA317/pLASN was the first therapeutic vector used in a gene therapy trial. (Blaese et al., Science 270:475-480 (1995)). Transduction efficiencies of 50% or greater have been observed for MFG-S packaged vectors. (Ellem et al., Immunol Immunother. 44(1): 10-20 (1997); Dranoff et al., Hum. Gene Ther. 1:111-2 (1997).

[0153] Recombinant adeno-associated virus vectors (rAAV) are a promising alternative gene delivery systems based on the defective and nonpathogenic parvovirus adeno-associated type 2 virus. All vectors are derived from a plasmid that retains only the AAV 145 bp inverted terminal repeats flanking the transgene expression cassette. Efficient gene transfer and stable transgene delivery due to integration into the genomes of the transduced cell are key features for this vector system. (Wagner et al., Lancet 351:9117 1702-3 (1998), Kearns et al., Gene Ther. 9:748-55 (1996)). Other AAV serotypes, including AAV1, AAV2, AAV3, AAV4, AAV5, AAV6, AAV7, AAV8, AAV9 and AAVrh.10 and any novel AAV serotype can also be used in accordance with the present invention.

[0154] Replication-deficient recombinant adenoviral vectors (Ad) can be produced at high titer and readily infect a number of different cell types. Most adenovirus vectors are engineered such that a transgene replaces the Ad E1a, E1b, or E3 genes; subsequently the replication defective vector is propagated in human 293 cells that supply deleted gene function in *trans*. Ad vectors can transduce multiple types of tissues *in vivo*, including nondividing, differentiated cells such as those found in liver, kidney and muscle. Conventional Ad vectors have a large carrying capacity. An example of the use of an Ad vector in a clinical trial involved polynucleotide therapy for antitumor immunization with intramuscular injection (Stermann et al., Hum. Gene Ther. 7:1083-9 (1998)). Additional examples of the use of adenovirus vectors for gene transfer in clinical trials include Rosenecker et al., Infection 24:1 5-10 (1996); Stermann et al., Hum. Gene Ther. 9:7 1083-1089 (1998); Welsh et al., Hum. Gene Ther. 2:205-18 (1995); Alvarez et al., Hum. Gene Ther. 5:597-613 (1997); Topf et al., Gene Ther. 5:507-513 (1998); Stermann et al., Hum. Gene Ther. 7:1083-1089 (1998).

[0155] Packaging cells are used to form virus particles that are capable of infecting a host cell. Such cells include HEK293 and Sf9 cells, which can be used to package AAV and adenovirus, and ψ 2 cells or PA317 cells, which package retrovirus. Viral vectors used in gene therapy are usually generated by a producer cell line that packages a nucleic acid vector into a viral particle. The vectors typically contain the minimal viral sequences required for packaging and subsequent integration into a host (if applicable), other viral sequences being replaced by an expression cassette encoding the protein to be expressed. The missing viral functions are supplied in *trans* by the packaging cell line. For example, AAV vectors used in gene therapy typically only possess inverted terminal repeat (ITR) sequences from the AAV genome which are required for packaging and integration into the host genome. Viral DNA is packaged in a cell line, which contains a helper plasmid encoding the other AAV genes, namely *rep* and *cap*, but lacking ITR sequences. The cell line is also infected with adenovirus as a helper. The helper virus promotes replication of the AAV vector and expression of AAV genes from the helper plasmid. The helper plasmid is not packaged in significant amounts due to a lack of ITR

sequences. Contamination with adenovirus can be reduced by, e.g., heat treatment to which adenovirus is more sensitive than AAV. In some embodiments, AAV is produced using a baculovirus expression system (see e.g. U.S. Patent Nos. 6,723,551 and 7,271,002).

[0156] Purification of AAV particles from a 293 or baculovirus system typically involves growth of the cells which produce the virus, followed by collection of the viral particles from the cell supernatant or lysing the cells and collecting the virus from the crude lysate. AAV is then purified by methods known in the art including ion exchange chromatography (e.g. see U.S. Patent Nos. 7,419,817 and 6,989,264), ion exchange chromatography and CsCl density centrifugation (e.g. PCT publication WO2011094198A10), immunoaffinity chromatography (e.g. WO2016128408) or purification using AVB Sepharose (e.g. GE Healthcare Life Sciences).

[0157] In many gene therapy applications, it is desirable that the gene therapy vector be delivered with a high degree of specificity to a particular tissue type. Accordingly, a viral vector can be modified to have specificity for a given cell type by expressing a ligand as a fusion protein with a viral coat protein on the outer surface of the virus. The ligand is chosen to have affinity for a receptor known to be present on the cell type of interest. For example, Han et al., Proc. Natl. Acad. Sci. USA 92:9747-9751 (1995), reported that Moloney murine leukemia virus can be modified to express human heregulin fused to gp70, and the recombinant virus infects certain human breast cancer cells expressing human epidermal growth factor receptor. This principle can be extended to other virus-target cell pairs, in which the target cell expresses a receptor and the virus expresses a fusion protein comprising a ligand for the cell-surface receptor. For example, filamentous phage can be engineered to display antibody fragments (e.g., Fab or Fv) having specific binding affinity for virtually any chosen cellular receptor. Although the above description applies primarily to viral vectors, the same principles can be applied to nonviral vectors. Such vectors can be engineered to contain specific uptake sequences which favor uptake by specific target cells.

[0158] Gene therapy vectors can be delivered *in vivo* by administration to an individual patient, typically by systemic administration (e.g., intravenous, intraperitoneal, intramuscular, subdermal, or intracranial infusion) or topical application, as described below. Alternatively, vectors can be delivered to cells *ex vivo*, such as cells explanted from an individual patient (e.g., lymphocytes, bone marrow aspirates, tissue biopsy) or universal donor hematopoietic stem cells, followed by reimplantation of the cells into a patient, usually after selection for cells which have incorporated the vector.

[0159] Vectors (e.g., retroviruses, adenoviruses, liposomes, etc.) containing nucleases, donor constructs, and combinations of nucleases and donor constructs can also be administered directly to an organism for transduction of cells *in vivo*. Alternatively, naked DNA can be administered. Administration is by any of the routes normally used for introducing a molecule into ultimate contact with blood or tissue cells including, but not limited to, injection, infusion, topical application and electroporation. Suitable methods of administering such nucleic acids are available and well known to those of skill in the art, and, although more than one route can be used to administer a particular composition, a particular route can often provide a more immediate and more effective reaction than another route.

[0160] Vectors suitable for introduction of polynucleotides (e.g. nuclease-encoding, double-stranded donors, and combinations of nuclease-encoding and double-stranded donors) described herein include non-integrating lentivirus vectors (IDLV). See, for example, Ory et al. (1996) Proc. Natl. Acad. Sci. USA 93:11382-11388; Dull et al. (1998) J. Virol. 72:8463-8471; Zuffery et al. (1998) J. Virol. 72:9873-9880; Follenzi et al. (2000) Nature Genetics 25:217-222; U.S. Patent Publication No. 2009/0117617.

[0161] Pharmaceutically acceptable carriers are determined in part by the particular composition being administered, as well as by the particular method used to administer the composition. Accordingly, there is a wide variety of suitable formulations of pharmaceutical compositions available, as described below (see, e.g.,

Remington's Pharmaceutical Sciences, 17th ed., 1989).

[0162] It will be apparent that the nuclease-encoding sequences and donor constructs can be delivered using the same or different systems. For example, the nucleases and donors can be carried by the same DNA MC. Alternatively, a donor polynucleotide can be carried by a MC, while the one or more nucleases can be carried by a standard plasmid or AAV vector. Furthermore, the different vectors can be administered by the same or different routes (intramuscular injection, tail vein injection, other intravenous injection, intraperitoneal administration or intramuscular injection). The vectors can be delivered simultaneously or in any sequential order.

[0163] Thus, the instant disclosure includes *in vivo* or *ex vivo* treatment of diseases and conditions that are amenable to insertion of a transgene encoding a therapeutic protein. The compositions are administered to a human patient in an amount effective to obtain the desired concentration of the therapeutic polypeptide in the serum or the target organ or cells. Administration can be by any means in which the polynucleotides are delivered to the desired target cells. For example, both *in vivo* and *ex vivo* methods are contemplated. Intravenous injection to the portal vein is a preferred method of administration. Other *in vivo* administration modes include, for example, direct injection into the lobes of the liver or the biliary duct and intravenous injection distal to the liver, including through the hepatic artery, direct injection in to the liver parenchyma, injection via the hepatic artery, and retrograde injection through the biliary tree. *Ex vivo* modes of administration include transduction *in vitro* of resected hepatocytes or other cells of the liver, followed by infusion of the transduced, resected hepatocytes back into the portal vasculature, liver parenchyma or biliary tree of the human patient, see e.g., Grossman et al., (1994) *Nature Genetics*, 6:335-341.

[0164] The effective amount of nuclease(s) and donor to be administered will vary from patient to patient and according to the therapeutic polypeptide of interest. Accordingly, effective amounts are best determined by the physician administering the compositions and appropriate dosages can be determined readily by one of ordinary skill in the art. After allowing sufficient time for integration and expression (typically 4-15 days, for example), analysis of the serum or other tissue levels of the therapeutic polypeptide and comparison to the initial level prior to administration will determine whether the amount being administered is too low, within the right range or too high. Suitable regimes for initial and subsequent administrations are also variable, but are typified by an initial administration followed by subsequent administrations if necessary. Subsequent administrations may be administered at variable intervals, ranging from daily to annually to every several years. One of skill in the art will appreciate that appropriate immunosuppressive techniques may be recommended to avoid inhibition or blockage of transduction by immunosuppression of the delivery vectors, see e.g., Vilquin et al., (1995) *Human Gene Ther.*, 6:1391-1401.

[0165] Formulations for both *ex vivo* and *in vivo* administrations include suspensions in liquid or emulsified liquids. The active ingredients often are mixed with excipients which are pharmaceutically acceptable and compatible with the active ingredient. Suitable excipients include, for example, water, saline, dextrose, glycerol, ethanol or the like, and combinations thereof. In addition, the composition may contain minor amounts of auxiliary substances, such as, wetting or emulsifying agents, pH buffering agents, stabilizing agents or other reagents that enhance the effectiveness of the pharmaceutical composition.

Cells

[0166] Also described herein are cells and cell lines in which an endogenous BCL11A enhancer sequence is modified by the nucleases described herein (Table 1). The modification may be, for example, as compared to the wild-type sequence of the cell. The cell or cell lines may be heterozygous or homozygous for the modification. The modifications to the BCL11A sequence may comprise indels.

[0167] The modification is preferably at or near the nuclease(s) binding site(s), cleavage site(s) and combinations of binding site(s) and cleavage site(s), for example, within 1-300 (or any value therebetween) base pairs upstream or downstream of the site(s) of cleavage, more preferably within 1-100 base pairs (or any value therebetween) of either side of the binding site(s), cleavage site(s), or binding site(s) and cleavage site(s), even more preferably within 1 to 50 base pairs (or any value therebetween) on either side of the binding site(s), cleavage site(s), or binding site(s) and cleavage site(s). In certain embodiments, the modification is at or near the "+58" region of the BCL11A enhancer, for example, at or near a nuclease binding site shown in any of the first column of Table 1.

[0168] Any cell or cell line may be modified, for example a stem cell, for example an embryonic stem cell, an induced pluripotent stem cell, a hematopoietic stem cell, a neuronal stem cell and a mesenchymal stem cell. Other non-limiting examples of cells as described herein include T-cells (e.g., CD4+, CD3+, CD8+, etc.); dendritic cells; B-cells. A descendent of a stem cell, including a partially or fully differentiated cell, is also provided (e.g., a RBC or RBC precursor cell). Non-limiting examples other cell lines including a modified BCL11A sequence include COS, CHO (e.g., CHO-S, CHO-K1, CHO-DG44, CHO-DUXB11, CHO-DUKX, CHOK1SV), VERO, MDCK, WI38, V79, B14AF28-G3, BHK, HaK, NS0, SP2/0-Ag14, HeLa, HEK293 (e.g., HEK293-F, HEK293-H, HEK293-T), and perC6 cells as well as insect cells such as *Spodoptera frugiperda* (Sf), or fungal cells such as *Saccharomyces*, *Pichia* and *Schizosaccharomyces*.

[0169] The cells as described herein are useful in treating or preventing a disorder, for example, by *ex vivo* therapies. The nuclease-modified cells can be expanded and then reintroduced into the patient using standard techniques. See, e.g., Tebas et al. (2014) New Eng J Med 370(10):901. In the case of stem cells, after infusion into the subject, *in vivo* differentiation of these precursors into cells expressing the functional transgene also occurs. Pharmaceutical compositions comprising the cells as described herein are also described. In addition, the cells may be cryopreserved prior to administration to a patient.

[0170] Any of the modified cells or cell lines disclosed herein may show increased expression of gamma globin. Compositions such as pharmaceutical compositions comprising the genetically modified cells as described herein are also described.

Applications

[0171] The methods and compositions disclosed herein are for modifying expression of protein, or correcting an aberrant gene sequence that encodes a protein expressed in a genetic disease, such as a sickle cell disease or a thalassemia. Thus, the methods and compositions provide for the treatment or prevention of such genetic diseases. Genome editing, for example of stem cells, can be used to correct an aberrant gene, insert a wild type gene, or change the expression of an endogenous gene. By way of non-limiting example, a wild type gene, e.g. encoding at least one globin (e.g., α globin, γ globin, β globin and combinations thereof), may be inserted into a cell (e.g., into an endogenous BCL11A enhancer sequence using one or more nucleases as described herein) to provide the globin proteins deficient or lacking in the cell and thereby treat a genetic disease, e.g., a hemoglobinopathy, caused by faulty globin expression. Alternatively or in addition, genomic editing with or without administration of the appropriate donor, can correct the faulty endogenous gene, e.g., correcting the point mutation in α - or β -hemoglobin, to restore expression of the gene or treat a genetic disease, e.g. sickle cell disease, knock out or alteration (overexpression or repression) of any direct or indirect globin regulatory gene (e.g. inactivation of the γ globin-regulating gene BCL11A or the BCL11A-regulator KLF1). Specifically, the compositions of the invention have use in the treatment or prevention of hemoglobinopathies.

[0172] The nucleases of the invention are targeted to the BCL11A enhancer region, known to be required for the expression of BCL11A during erythropoiesis, and hence the down regulation of gamma globin expression.

Modification of this enhancer region may result in erythrocytes with increased gamma globin expression, and thus may be helpful for the treatment or prevention of sickle cell disease or beta thalassemia.

[0173] The following Examples relate to exemplary embodiments of the present disclosure in which the nuclease comprises a zinc finger nuclease (ZFN). It will be appreciated that this is for purposes of exemplification only and that other nucleases can be used, for example TtAgo and CRISPR/Cas systems, homing endonucleases (meganucleases) with engineered DNA-binding domains, fusions of naturally occurring of engineered homing endonucleases (meganucleases) DNA-binding domains, including combinations of homing endonucleases (meganucleases) with engineered DNA-binding domains and fusions of naturally occurring of engineered homing endonucleases (meganucleases) DNA-binding domains, and heterologous cleavage domains, fusions of meganucleases and TALE proteins, including combinations of heterologous cleavage domains and fusions of meganucleases and TALE proteins.

EXAMPLES

Example 1: Assembly of Zinc Finger Nucleases

[0174] ZFNs were assembled against the human BCL11A gene and activity was tested by deep sequencing analysis of DNA isolated from transfected cells as described below. ZFNs specific for the +58 region of the enhancer region were made as described. ZFN pair 51857/51949 has been described previously (see WO 2016/183298).

Example 2: Off target analysis

[0175] To analyze off target cleavage by the ZFN pairs, a two stage unbiased specificity analysis was performed. In the first stage, (Figure 1), candidate off-target sites for each ZFN were identified via an oligonucleotide duplex integration site assay using a procedure similar to that described by Tsai et al. ((2015), Nat Biotechnol 33(2): 187-197. doi: 10.1038/nbt.3117).

[0176] The oligonucleotide duplex integration site assay is based on the observation that co-introduction of a nuclease and a short segment of duplex DNA into a target cell results in duplex integration during repair of a fraction of genome cleavage events via the NHEJ DNA repair pathway (Orlando et al., (2010), Nucleic Acids Res, 38(15) e152. doi: 10.1093/nar/gkq512; Gabriel et al., (2011), Nat Biotechnol. 2011 Aug 7;29(9):816-23. doi: 10.1038/nbt. 1948; Tsai *et al.*, *ibid*). Upon integration the duplex provides a permanent tag of the cleavage event. Sites of integration are then identified via ligation of an oligonucleotide adaptor to sheared genomic DNA, followed by 2 rounds of 25 cycles of nested PCR, and deep sequencing of the resulting donor-genome junctions. This assay allows for evaluation of all potential integration sites within the genome.

[0177] The integration site assay was performed in K562 cells to maximize donor delivery, ZFN expression, and donor integration. Moreover as K562 cells divide quickly (doubling time approximately 24 hours) they are expected to impose minimal epigenetic restrictions on the ability of ZFNs to cleave cellular targets. Cells (2×10^5) were electroporated with 0.47 μ g of oligonucleotide duplex donor and 400 ng of each ZFN-encoding mRNA using an Amaxa shuttle and settings optimized for maximal on-target activity of the ZFNs. Four replicate samples were prepared for each combination of oligo and mRNA. On day 7 post-transfection, genomic DNA was isolated for each sample (Qiagen DNeasy Blood and Tissue Kit) and 400 ng (133000 haploid genomes) was used as input for the amplification protocol outlined in Figure 1. Samples were then processed essentially as described (Tsai *et al.* *ibid*). Final products were pooled, quantified, and sequenced on

a MiSeq Instrument (Illumina) using a v2 300 cycle sequencing kit with paired-end 150 bp reads and 8 bp/16 bp dual index reads to detect the sample barcodes on each end of the amplicon.

[0178] To generate a list of candidate off target sites, sequencing data were filtered for correct priming sequence, followed by trimming of adapter sequences and mapping to the genome. Next, junction coordinates were mapped, and the duplex-genome junction, as well as the position of the break caused by DNA shearing, were used to identify distinct integration events. Integration events were then processed to identify clusters of integrations in close proximity within the genome (minimum of 4 distinct integration events within 100bp of each other, summed across all replicates). Clusters residing on contigs that were unmappable in the hg38 assembly (*i.e.* chrUn in hg38) were removed from further analysis. Clusters mapping to repetitive loci (median of three or more hits to the genome across all sequences in a cluster) were also removed as prior experience has shown these to be amplification artifacts. Remaining clusters were scored as candidate ZFN cleavage sites if they were derived from at least 2 replicate ZFN treated samples (of 4 total) and exhibited ≥ 5 fold excess of integration events in ZFN treated samples versus controls. Candidate cleavage sites were ranked by the total number of unique integrations in the ZFN treated samples. Candidate loci identified via this analysis are provided in Figure 2 for ZFN pair 51857/51949, ranked by integrant count.

Example 3: Optimization of ZFNs

[0179] To decrease off target cleavage, a strategy for nuclease optimization in which nonspecific phosphate contacts are selectively removed to bring about global suppression off-target cleavage (Guillinger et al. (2014) Nat Methods. 11(4):429-35. doi: 10.1038/nmeth.2845; Kleinstiver et al. (2016) Nature 529(7587):490-5. doi: 10.1038/nature16526; Slaymaker et al. (2016) Science) 351(6268):84-8. doi: 10.1126/science.aad5227) was adopted (see U.S. Provisional Application Nos. 62/443,981 and 62/378,978). Amino acid substitutions were made at a key position within the zinc finger framework that interacts with the phosphate backbone of the DNA (Pavletich and Pabo, (1991) Science 252(5007):809-17; Elrod-Erickson et al. (1996) Structure 4(10): 1171-80) (Figure 3A-3B) as well as at a single position in the right ZFN FokI domain also predicted to make a phosphate contact (Figure 3C).

[0180] Specificity was further improved by allowing independent expression of each ZFN from two separated mRNAs, which enables optimization of delivery ratios. These efforts yielded optimized ZFN pairs that are highly related to the original one, differing by substitutions that decrease the energetics of interaction with the DNA phosphate backbone but that minimally or do not impact sequence specific base recognition. Consistent with this, the integration site assay yielded 455 loci for potential targets of ZFN cleavage for the original 51857/51949 pair. For the optimized pair, a much smaller number of loci were identified for further examination as potential targets of ZFN cleavage (72 total) by this analysis. For both pairs, the intended target within the BCL11A enhancer was the top ranked locus. Moreover a much higher fraction of integration events was noted at the BCL11A enhancer for the optimized pair, consistent with its greater specificity.

[0181] It is important to note that in defining the sequence data processing pipeline, key parameters were chosen conservatively, to err on the side of including as many candidate off target loci as feasible instead of filtering them out. This was done to ensure that every locus that might represent a bona fide cleavage site for the optimized ZFNs would be identified and tested in follow-up indel studies, even at the cost of accepting a much greater number that would become false positives. It was expected that the first stage of analysis would yield a large set of candidate loci for each ZFN pair, of which the large majority (particularly for the optimized ZFNs) would not represent true off-target cleavage sites but rather background events that would prove negative for cleavage in follow-up indel studies.

[0182] In the second stage of analysis, candidate off-target loci identified via the integration site assay were screened for evidence of modification (*e.g.*, the presence of indels) in ZFN-treated CD34+ HSPC.

[0183] In particular, human CD34+ HSPC derived from mobilized peripheral blood were treated with the original and optimized ZFN pairs using clinical scale and clinical conditions for RNA transfection (120 µg/mL of mRNA for the original ZFN pair and 100 µg/mL of mRNA for the optimized pair). Genomic DNA was isolated 2 days post-transfection, followed by PCR amplification of candidate off-target loci and deep sequencing to quantify indel levels. For both the original and optimized ZFN pairs, the same set of 137 candidate off-target loci were screened at this step, along with a smaller number of candidate off-target sites that had been identified via other methods in earlier studies with the original ZFNs.

[0184] The results showed that the optimized ZFNs are markedly more specific than the original pair. This is apparent not only from the number of loci that were scored positive for evidence of ZFN cleavage (52 for the original pair vs 3 for the optimized pair), but also from the observed indel levels, which for the optimized pair were much lower. Figure 4 shows plots of indel values at every locus exhibiting evidence of ZFN cleavage in this study (note log scale of y axis). Aggregating off-target indels across all such loci indicates a reduction in off-target activity of 300 fold (46.5% aggregate off-target indels for the original pair, vs 0.15% off-target indels for the optimized pair). This reduction in off-target activity was achieved without any loss in activity at the intended target site (72.5% indels for the original pair vs 81.9% for the optimized ZFNs). In these studies, the original pair (or parental pair) was 51857/51949, while the optimized ZFN pair was 63014/65722 (see below).

[0185] The nuclease designs are shown below in Table 1:

Table 1: ZFN pairs specific for +58 BCL11A enhancer region

SBS # (target site, 5'-3')	Design [Helix Sequence, SEQ ID]						Linker
	[Mutations to finger backbone]						Fok mutants
	F1	F2	F3	F4	F5	F6	
Left partner							
51857 aaAGCAACTg TTAGCTTGCA Ctagacta (SEQ ID NO:1)	DQSNLRA (SEQ ID NO:19)	RNFSLTM (SEQ ID NO:20)	STGNLTN (SEQ ID NO:21)	TSGSLTR (SEQ ID NO:22)	DQSNLRA (SEQ ID NO:19)	AQCCLFH (SEQ ID NO:23)	L7c5
	none	none	none	none	none	none	ELD
63014 aaAGCAACTg TTAGCTTGCA Ctagacta (SEQ ID NO:1)	DQSNLRA (SEQ ID NO:19)	RNFSLTM (SEQ ID NO:20)	STGNLTN (SEQ ID NO:21)	TSGSLTR (SEQ ID NO:22)	DQSNLRA (SEQ ID NO:19)	AQCCLFH (SEQ ID NO:23)	L7c5
	Qm5	none	Qm5	none	Qm5	none	ELD
65459 aaAGCAACTg TTAGCTTGCA Ctagacta (SEQ ID NO:1)	DQSNLRA (SEQ ID NO:19)	RNFSLTM (SEQ ID NO:20)	STGNLTN (SEQ ID NO:21)	TSGSLTR (SEQ ID NO:22)	DQSNLRA (SEQ ID NO:19)	AQCCLFH (SEQ ID NO:23)	L7c5
	Qm14Qm5	none	Qm5	none	Qm5	none	ELD
Right Partner							
51949 caCAGGCTCC AGGAAGGgtt tggcctct (SEQ ID NO:2)	RNDHRTT (SEQ ID NO:24)	QKAHLIR (SEQ ID NO:25)	QKGTLGE (SEQ ID NO:26)	RGRDLSR (SEQ ID NO:26)	RRDNLHS (SEQ ID NO:27)	N/A	LO
	none	none	none	none	none	N/A	KKR
65722 caCAGGCTCC AGGAAGGgtt tggcctct (SEQ ID NO:2)	RNDHRTT (SEQ ID NO:24)	QKAHLIR (SEQ ID NO:25)	QKGTLGE (SEQ ID NO:26)	RGRDLSR (SEQ ID NO:26)	RRDNLHS (SEQ ID NO:27)	N/A	LO
	Qm5	Qm5	none	Qm5	none	N/A	KKR K525S

SBS # (target site, 5'-3')	Design [Helix Sequence, SEQ ID]						Linker
	[Mutations to finger backbone]						Fok mutants
	F1	F2	F3	F4	F5	F6	
Left partner							
65526 caCAGGCTCC AGGAAGGgtt tggcctct (SEQ ID NO:2)	RNDHRTT (SEQ ID NO:24)	QKAHLIR (SEQ ID NO:25)	QKGTIGE (SEQ ID NO:26)	RGRDLSR (SEQ ID NO:26)	RRDNLHS (SEQ ID NO:27)	N/A	LO
	Qm5	Qm5	none	Qm5	Qm5	N/A	KKR R416S
65549 caCAGGCTCC AGGAAGGgtt tggcctct (SEQ ID NO:2)	RNDHRTT (SEQ ID NO:24)	QKAHLIR (SEQ ID NO:25)	QKGTIGE (SEQ ID NO:26)	RGRDLSR (SEQ ID NO:26)	RRDNLHS (SEQ ID NO:27)	N/A	LO
	Qm5	Qm5	Qm5	Qm5	none	N/A	KKR K525S
65550 caCAGGCTCC AGGAAGGgtt tggcctct (SEQ ID NO:2)	RNDHRTT (SEQ ID NO:24)	QKAHLIR (SEQ ID NO:25)	QKGTIGE (SEQ ID NO:26)	RGRDLSR (SEQ ID NO:26)	RRDNLHS (SEQ ID NO:27)	N/A	LO
	Qm5	Qm5	none	Qm5	Qm5	N/A	KKR K525S

[0186] Table 1 shows characterizing information pertaining to each ZFN. Starting from the left, the SBS number (e.g. 51857) is displayed with the DNA target that the ZFN binds to displayed below the SBS number. Next are shown the amino acid recognition helix designs for fingers 1-6 or 1-5 (subdivided column 2 of Table 1). Also shown in Table 1 under the appropriate helix designs are mutations made to the ZFP backbone sequences of the indicated finger, as described in U.S. Provisional Patent Application Nos. 62/378,978 and 62/443,981. In the notation used in Table 1, "Qm5" means that at position **minus 5** (relative to the helix which is numbered -1 to +6) of the indicated finger, the arginine at this position has been replaced with a glutamine (**Q**), while "Qm14" means that the arginine (**R**) normally present in position **minus 14** has been replaced with a glutamine (**Q**). "None" indicates no changes outside the recognition helix region. Thus, for example, SBS# 63014 includes the Qm5 mutation in fingers 1, 3 and 5 while fingers 2, 4 and 6 do not have mutations to the zinc finger backbone (e.g., the zinc finger sequence outside the recognition helix region).

[0187] Finally, the right-most column of Table 1 shows the linker used to link the DNA binding domain to the FokI cleavage domain (e.g., "L7c5" (LRGSISRARPLNPHP (SEQ ID NO:5), as described for example in U.S. Patent No. 9,567,609) is displayed on top line of the column, with the sites of the FokI phosphate contact mutations and dimerization mutations shown in the box below the linker designation. In specifics, indicated on top line of the Fok mutants box is the type of mutation found in the dimerizing domain (e.g., ELD or KKR as described for example in U.S. Patent No. 8,962,281). Below the dimerization mutant designations is shown any mutations present in the FokI domain made to remove a non-specific phosphate contact shown on the bottom (e.g. K525S or R416S where serine residues at amino acid positions 525 or 416 have been substituted for either a lysine or arginine, respectively as described in U.S. Provisional Patent Application Nos. 62/378,978 and 62/443,981). Thus, for example, in SBS# 63014, the linker is an L7c5 linker and the FokI cleavage domain includes the ELD dimerization mutants and no phosphate contact mutations. Further, for SBS# 65722, the linker is an L0 linker (LRGSQLVKS (SEQ ID NO:6), also referred to as the 'standard' linker, see U.S. Patent No. 9,567,609) and the FokI cleavage domain includes the KKR dimerization mutations and the K525S FokI phosphate contact mutation.

[0188] All ZFNs were tested for functionality (cleavage activity as determined by assaying for indels as

described in Example 4 below) and found to be active.

[0189] Furthermore, in order to determine which ZFN designs were the most specific, indel analyses of known sites of off-target cleavage by the original ZFN pair were performed in ZFN-treated CD34⁺ HSPC. To accomplish this, human CD34⁺ HSPC derived from mobilized peripheral blood were treated with the original and optimized ZFN pairs using clinical conditions and mRNA concentrations (120 µg/mL for the original ZFN pair and 100 µg/mL for the optimized pair). Genomic DNA was isolated 2 days post-transfection from these cells and untreated controls, followed by PCR amplification of each candidate locus and deep sequencing to quantify indel levels.

[0190] Modification levels at each locus were determined by paired-end deep sequencing on an Illumina MiSeq using a 300 cycle cartridge. Paired sequences were merged, adaptor trimmed via SeqPrep filtered for a quality score of ≥ 15 across all bases, and then mapped to the human genome (hg38 assembly). Sequences that mapped to an incorrect locus were discarded. Sequences shorter than the wild-type amplicon by >70 bp or $>70\%$ were removed in order to minimize primer-dimer products. A Needleman-Wunsch alignment (Needleman and Wunsch, (1970), J Mol Biol 48(3):443-53)) was performed between the target amplicon and each MiSeq read to map indels. Indels in aligned sequences were defined as described in Gabriel *et al.* 2011 (*ibid*) except that indels 1bp in length were also accepted to avoid undercounting real events. Note that a fraction of loci either did not amplify or did not sequence, or were rejected from analysis due to high background ($>1\%$ modification in control samples) or insufficient sequencing depth (<10000 reads). The results of this analysis and comparison to the 'parent' 51857/51949 ZFN pair are provided below in Table 2.

Table 2: Off target cleavage analysis

ZFN dimer				Off-target: # of loci			# indel-positive OT loci		
Left	Right			target	PCR'd	Analyzable	P<0.05	manual	In parent? Capture/confirmed
51857	51949	60:60	73.0	23	21	17	15	17	-
63014	65722	60:15	82.2	31	24	14	0	4	4/3
63014	65526	60:15	81.2	23	22	9	3	4	4/4
63014	65527	60:60	81.4	30	24	10	4	4	4/2
63014	65549	60:60	80.0	30	24	13	0	2	2/1
63014	65550	60:60	79.8	30	24	9	0	1	1/1
65459	65526	60:15	76.9	23	19	14	0	2	2/2

Example 4: Activity of ZFNs in human CD34⁺ cells

[0191] For *in vitro* testing, the nucleases were tested in CD34⁺ cells. ZFNs were supplied as mRNAs, where the mRNAs were made *in vitro* as follows: plasmids comprising the genes encoding the ZFN are linearized and used for *in vitro* mRNA transcription using the mMessage mMachine® T7 Ultra Kit (Ambion/Applied Biosystems). The mRNA was then purified using an RNeasy® mini kit (Qiagen).

[0192] CD34⁺ cells were isolated from mobilized peripheral blood and maintained in X-VIVO 10 medium supplemented with penicillin, streptomycin and glutamine as well as StemSpan CC110 and incubated at 37°C and 5% CO₂. Cells were transfected 48 hours post-isolation or post-thaw. A small aliquot was mixed 1:1 with trypan blue solution 0.4% (w/v) in PBS (Corning) and the cell numbers were determined on a TC20 Automated Cell Counter (Bio-Rad).

[0193] For large scale transfections, cells were washed with MaxCyte Electroporation Buffer (Maxcyte) and re-suspended at 3 to 5e7 cells per mL in Electroporation buffer in 100 μ L. Typically, mRNA concentrations between 60 μ g/mL and 120 μ g/mL were used to screen candidate ZFN sets. Cells were then grown in growth media at 3e6 cells per mL for 18 hours at 30°C and then diluted to 1e6 cells per mL for an additional 24 hours at 37°C. For determination of cleavage activity, genomic DNA was isolated 2-3 days post-transfection, and the level of gene modification at the BCL11A enhancer locus was measured via deep sequencing on a MiSeq sequencer (Illumina).

[0194] The ZFN pairs from Table 1 were tested in CD34+ cells and the activity results are shown below in Table 3.

Table 3: Activity of ZFN pairs against BCL11A target

Right ZFN	Left ZFN	R ZFN conc. (μ g)	L ZFN conc. (μ g)	Indels (%)
51857	51949	60	60	72.98
63014	65722	15	60	80.62
63014	65722	60	15	82.19
63014	65526	60	15	81.22
63014	65527	60	60	81.43
63014	65549	60	60	79.82
63014	65550	60	60	79.96
GFP control				0.07

[0195] In addition to analyzing the nuclease activity in CD34+ cells prior to erythroid differentiation, edited cells were also differentiated *in vitro* into erythroid cells. The protocol followed was based on Giarratana et al., ((2011) Blood 120 (15):2945-53). In brief, the protocol below was followed:

[0196] Day 0 to Day 7: 4×10^4 CD34+ cells were cultured at a density of 2×10^4 /mL in differentiation medium (EDM) (Iscove's Modified Dulbecco's Medium [IMDM], 330 μ g/mL Transferrin ,10 μ g/mL Human Insulin, 2 U/mL Heparin sodium, 5% Human AB+ plasma) in the presence of 10^{-6} M hydrocortisone, 100 ng/mL stem cell factor (SCF), 5 ng/mL IL 3, and 3 IU/mL erythropoietin (EPO).

[0197] Day 4: Cells resuspended in fresh EDM containing SCF, IL-3, EPO, and hydrocortisone.

[0198] Day 7 to Day 11: Cells were resuspended at a density of 1.5×10^5 cells per fresh mL of EDM supplemented with SCF and EPO.

[0199] Day 11 to Day 21: On day 11, cells were replated at 1×10^6 /mL in fresh EDM supplemented with EPO. Cells were subsequently replated in this same media at 5×10^6 /mL on day 14. Growth plateaus during this period of time between day 14 to 18, when cell viabilities began to drop until termination of the cultures at day 21.

[0200] Cell counts were taken at the time of seeding and throughout the differentiation by measuring Acridine Orange positivity and Propidium Iodide exclusion (AOPI) using a Nexcelom Bioscience Cellometer K2 with the AOPI Erythroid Assay mode with fluorescence channel 1 (AO) set to 700 milliseconds and fluorescence channel 2 (PI) set to 5000 milliseconds.

[0201] The percentage of enucleated cells was determined at day 21 of the differentiation using the following protocol. The enucleation rate was comparable among untransfected controls and ZFN-transfected samples

with percentages 59-63% from these two groups:

1. 1. Cell count
2. 2. 100,000 cells, spin down at 450 x g, 5 min, RT.
3. 3. Resuspend in 50 μ L PBS-BSA + 1 μ L GlyA-FITC (DAKO).
4. 4. Stain for 15 min in fridge.
5. 5. Add 1 mL PBS-BSA, vortex, spin down.
6. 6. Resuspend in 250 μ L of PBS-BSA-NucRed (2 drops NucRed per mL).
7. 7. Acquire on FACS Canto using the APC channel for NucRed.
8. 8. Nucleated erythroid cells will be in the GlyA positive NucRed negative/low fraction and erythroblasts will be in the double GlyA-NucRed positive fraction.

[0202] BCL11A gene modification was measured by MiSeq deep sequencing in DNA samples harvested a) 48 hours after electroporation b) on the day of thawing the cells, at the time when the *in vitro* differentiation was started and c) at day 14 of the *in vitro* erythroid differentiation. While the differentiation was performed for 21 days, the day 14 timepoint for DNA analysis was chosen since it is prior to enucleation of a large fraction of the erythroid cells which results in a loss of DNA recovery. The observed modification percentages at the BCL11A enhancer are listed in Table 4 together with details of the transfection conditions.

Table 4: BCL11A Gene Modification Levels by MiSeq Analysis

CD34+ Cell Prep	Transfection	BCL11A Gene Modification (%)		
		Day 2 post TF	Post-Thaw	Day 14 of differentiation
Prep #1	80 μ g/mL 63014 + 20 μ g/mL 65722	78.6	81.4	72.0
	Untransfected	0.1	0.2	0.2
Prep#2	80 μ g/mL 63014 + 20 μ g/mL 65722	75.4	77.3	72.3
	Untransfected	0.1	0.1	0.0

[0203] These data show that CD34+ cell transfection with optimized pair 63014 and 65722 mRNA leads to very efficient gene modification at the BCL11A enhancer target site (>75% of alleles modified) and that the modification is maintained very well (>90% retention of the modification) after freezing and thawing of the cells and after erythroid differentiation.

[0204] The ZFN pair 63014/65722 was selected for further analysis. The amino acid sequences for these ZFNs are shown below, where each comprises a nuclear localization signal (NLS, Kalderon et al. (1984) Cell 39 (3 Pt 2):499-509) and a hydrophilic peptide (Hopp et al. (1988) Nat Biotechnol 6:1204-10) which enhances on-target ZFN activity, both fused to the N-terminal coding sequence. Thus, the mRNA and amino acid sequences of the ZFNs are as follows:

63014 mRNA (1725 nt)

5'gggagacaagcuuugaauuacaagcuugcuuuuuugcagaagcucagaauaaacgcucaacuuggcagauagcgaauucgccaugacuaaagaccagacggugauuuuaaagaucagacauagcgaauuacaaggauacgaugacaagaugcccccagaagaagaggagcggcauccacggguacccgccgcuauggcugagaggccuuccagugucgaauucgaugcagaacuucagugaccaguccaaccugcgcccacauccgcacccacacggcgagaagccuuuugccugacauuuugggaggaaauuugcccgaacuucuccugaccagcauacaaagauacacacgggcagccaaaagccuuccagugucgaauucgaugcagaacuucaguuaccggcaaccugaccaaccacauccgcacccacacggcgagaagccuuuugccugacauuuugggaggaaauuugccac

cucccgucuccugacccgccauaccagauacacgcaccgcgcgccccgaucgccgaagccuuccaguguc
gaauucugcaugcagaacucagugaccagucacaacucgcgcgcccauaccgcacccacaccgcgcgagaagccu
uuugccugugacauuuugugggaggaauuuugccgcgccaguguuugucuguccaccuaccagauacaccucg
ggggauccaucagcagagccagaccacugaacccgcaccgagcugcgagggagaaggaugccgagcugcgcgac
aagcugagaugacugccccacagagauacucgagcugaucgagaucgccaggaacagcaccacgagaccgcauccu
ggagauagaagugaugggagauucucaugagauguaucgcuacaggggaaagcaccugggcggaagcagaaa
gccugacggcgccaucuaucagugggcgagcccaucgaauacggcgugaucgugacacaaaggccuacagc
ggcgcguaacaucugccuauccggccagggccgacgagauaggagagauacgugggaggaacacgacccgggaa
agcaccucaaccccaacgagugugugaauguguaaccuagcagcugacaggauucaaguuccugucgugag
cggccacuucaaagggaacuaacaaggcccgugaccaggcugaaaccauacaccaacugcaauggcgccgugc
ugagcgugggagggagcugcugaucggcgcgagaugaucaaagccggcacccugacacugagggagggugcgcg
gcaaguucaacaacggcggaucacauucagauucugauaacucgagucucagaagcucgcuuucucugucc
aaauucuaauaaaggguuccuuuguuuccuaaguccaacuacuaaacugggggauauuugaaggggccuugagc
aucuggaucucgccuauuaaaaaacauuuuuuucuuugcugcgcuagaagcucgcuuucucuguccaaau
ucuaauaaaggguuccuuuguuuccuaaguccaacuacuaaacugggggauauuugaaggggccuugagcauc
ggauucugccuauuaaaaaacauuuuuuucuuugcugcgggacauucuaauuaaaaaaaaaaaaaaaaaaa
aacuag

(SEQ ID NO:28).

[0205] 63014 amino acid sequence (recognition helix regions are underlined; linker is shown in upper case italics; mutations to fingers 1, 3 and 5 backbone residues are shown in double-underlining; the dimerization domain mutations (ELD) are shown in bold and italics; hydrophilic peptide is indicated in lower case text; and the nuclear localization signal (NLS) is shown in lowercase italics):

Mdykdhdgdgdykdhdidyd kddddd kMApkkkrkvGIHGVPAAMAERPFQCRICMQNFSDQS
NLRAHIRTHTGEKPFACDICGRKFARNFSLTMHTKIHTGSQKPFQCRICMQNF
SSTGNLTNHIRTHTGEKPFACDICGRKFATSGSLTRHTKIHTHPRAPIPKPFQCR
ICMQNFSDQSNLRAHIRTHTGEKPFACDICGRKFAAQCCLFHHTKIHLGRSISR
ARPLNPHELEKKSELRHKLKYVPHEYIELIEIARNSTQDRILEMKVMEFFMK
VYGYRGKHLGGRKPDGAITYVGSPIDYGVIVDTKAYSGGYNLPIGQADEME
RYVEENQTRDKHLNPNNEWVKVYPSSVTEFKFLFVSGHFKGNKYAQLTRLNH
ITNCNGAVLSVEELLIGGEMIKAGTLTLEEVRRKFNNGEINFRS (SEQ ID
NO:29).

[0206] 65722 mRNA (1680 nucleotides):

5' gggagacaagcucugaauacaaagcucugucuucuuuuuugagcagaagcucagaauaaacgcucauaauuuggca
gaucgaauucgccuagagaucugcgcgcgcgagagggcgagagggagagucuuaaccugcgcgugacguugagagga
gaaucccgcccuagcgaccauggacuacaagaccugacgugauuuaaaagaucaugacaucgaauacaaaggg
augacguagacaaagauggccccaagaagaaagaggaagugucggauucaugggguacccgccgcuauggcuga
gagggccuuuccagugucgaaucugcaugcagaaguugcccgcaacgaccaccgaccacccuaaacagaauc
acacggcgcgagaaagccuuuccagugucgaaucugcaugcagaacuucagucagaagggccaccugauccgcccac
auccgcacccacaccggcgagaagccuuuugccugugacauuugugggaggggaauuugccagaagggcgaccc
ugggcgagcuauacaagauacacacgggaucucagaagccuuuccagugucgaaucugcaugcagaacuucag
ucggcgccgcgaccugucccgccacauccgcacccacaccggcgagagggccuuuugccugugacauuuguggg
agggaauuugcccgccgcgacaacugcacuccauccaagaauacaccugcgcggaugccagcugugagaagag

[illegible]

[0207] 65722 amino acid sequence (recognition helix regions are underlined; linker is shown in upper case italics; hydrophilic peptide is in lower case; nuclease localization signal is in lower case italics; mutations to fingers 1, 2 and 4 backbone residues are shown in double-underlining; the dimerization domain mutations (ELD) are shown in bold and italics; and the FokI phosphate contact mutation(s) is shown in wavy underlining):
Mdykdhdgdykdhdidykd d d d d k M A *p k k k r k v* G I H G V P A A M A E R P F Q C R I C M Q K F A R N D
H R T T H T K I H T G E K P F Q C R I C M Q N F S Q K A H L I R H I R T H T G E K P F A C D I C G R K F A Q
K G T L G E H T K I H T G S Q K P F Q C R I C M Q N F S R G R D L S R H I R T H T G E K P F A C D I C G R
K F A R R D N L H S H T K I H L R G S Q L V K S E L E E K K S E L R H K L K Y V P H E Y I E L I E I A R N S T
Q D R I L E M K V M E F F M K V Y G Y R G K H L G G S R K P D G A I Y T V G S P I D Y G V I V D T K A
Y S G G Y N L P I G Q A D E M Q R Y V *K E N* Q T R N K H I N P N E W W K V Y P S S V T E F K F L F V S

G H F S G N Y K A Q L T R L N *R K* T N C N G A V L S V E E L L I G G E M I K A G T L T L E E V R R K F N
N G E I N F (S E Q I D N O : 31).

Example 5: Assessment of Globin levels in Erythroid Progeny

[0208] Levels of α -, β - and γ - globin mRNA in cellular mRNA isolated at day 14 of the differentiation (before overall mRNA levels decline dramatically in the course of enucleation and erythroid maturation) were determined by RT-qPCR for the two cell preps shown above in Table 4. The γ - globin mRNA values are shown normalized relative to the β - globin mRNA (Figure 5A) or to the α - globin mRNA values (Figure 5B) from the same samples (using arbitrary units based on the ratio in the untransfected RT-PCR standard defined as 1).

[0209] Reverse phase HPLC of protein samples isolated at day 21, the end point of the erythroid differentiation, was used to determine whether ZFN mediated modification of the BCL11A erythroid enhancer elevates fetal hemoglobin at the protein level. The Gamma globin (sum of the Agamma and Ggamma peaks) to alpha globin ratios were determined, as well as the Gamma globin (sum of the Agamma and Ggamma peaks)/ over beta-like globin ratios (sum of the Agamma, Ggamma, beta and delta- globin peaks) and are shown in Figure 6.

[0210] In this experiment an approximate 3-4 fold elevation of fetal globin protein percentages to levels of about 15%-20% was observed in erythroid progeny of HSPCs upon 63014/65722-mediated disruption of the

BCL11A enhancer.

Example 6: Engraftment of edited cells in NSG mice

[0211] Edited human CD34⁺ cells were then injected into NSG mice to assess engraftment. The extent of human chimerism (*i.e.* the percentage of human CD45⁺ cells) was measured using fluorescence activated cell sorting (FACS) in peripheral blood collected at 8, and 12, 16 and 20 weeks post transplantation, and in bone marrow collected at 12 weeks and 20 weeks. In addition, to test the level of engraftment of ZFN modified cells, the level of gene disruption at the BCL11A enhancer locus was evaluated by direct high-throughput sequencing of the ZFN target locus and compared to the levels of target gene modification measured in the input material.

[0212] HSPC from two healthy donors (termed PB-MR-003 and PB-MR-004) were mobilized with G-SCF and Plerixafor and purified as described in Yannaki et al. ((2012) Mol Ther 20(1):230-8. doi: 10.1038/mt.2011). Platelet depletion was performed on the leukapheresis product using the Fresenius-Kabi Lovo device before it was enriched for CD34⁺ cells using the Miltenyi Biotech CliniMACS Plus instrument. The purified cells were then seeded in culture for transfection.

[0213] Two days after CD34⁺ cell purification, the cells were electroporated using the Maxcyte instrument in the presence of either 120 µg/mL of a single mRNA encoding the parental ZFN pair, 63014/65722 or optimized amounts of two separate mRNAs encoding the optimized ZFN pair 80 µg/mL 63014 and 20 µg/mL 65722. Before transfection, an aliquot of the cells were set aside as the untransfected control. 95 million cells were transfected from PB-MR-003 and 120 million cells were transfected from PB-MR-004.

[0214] Following electroporation, a transient overnight culture at 30°C was performed and cells were then cultured for an additional 24 hours at 37°C. Two days post-electroporation, cells aliquots for DNA analysis were taken and the remaining cells were harvested, cryopreserved, and stored in liquid nitrogen.

[0215] Conditioning: Mice were treated with 10 mg/kg/day Baytril water 1-2 days prior to irradiation and sublethally irradiated with 300 RAD 16-24 hours before transplantation. Transplantation was performed via tail vein injection (see below). Then mice received fresh Baytril water. Baytril water was replace one week later and Baytril water addition was discontinued 14 days after transplantation.

[0216] Transplantation: On the day of transplantation, prewarm X-Vivo 10/ 1%PSG + 3 cytokine cocktail (Recombinant Human Stem Cell Factor (SCF), Recombinant Human Thrombopoietin (TPO), and Recombinant Human Flt-3 Ligand (Flt-3L)) at 37°C, prepare fresh PBS/0.1% BSA at ambient temperature (sterile/filtered). The cryopreserved cells were thawed at 37°C, pelleted, resuspended in pre-warmed X-Vivo medium, pelleted again, resuspended in PBS/0.1% BSA and counted. After another pelleting the cell pellet was resuspended in 550 µL per mouse of PBS/0.1%BSA (2 x 10⁶ cells/mL based off the cell count). Cells were then injected at room temperature into mouse tail vein with a 25 gauge needle. Study groups are shown below in Table 5.

Table 5: Dosing groups for Engraftment of edited hCD34⁺ cells

Species: Mouse				Sex, Age, Number: 60 Female NSG Mice			
Group No.	N/ Group	Test Article	% indels (Day 2)	Viability 1d Post-Thaw	Dose (cells/mouse)	N / Sacrifice	
						Week 12	Week 20
1	10 (M1-10)	PB-MR-003 donor cells, treated with 63014 and 65722 mRNA.	79%	83%	1 million	5	5
3	10 (M21-30)	PB-MR-003.untransfected	0.1%	95%	1 million	5	5
4	10 (M31-40)	PB-MR-004 donor cells treated with 63014 and 65722 mRNA.	75%	77%	1 million	5	5
6	10 (M51-60)	PB-MR-004.untransfected	0.1%	92%	1 million	5	5

[0217] Animals were observed daily for general health and weighed daily for the first 2 weeks and weighed bi-weekly thereafter. Peripheral blood was collected from the submandibular vein (100 µL) at 8, 12, 16 and 20 weeks post transplantation or via cardiac puncture (1 mL) for the sacrificed animals at 12 and 20 weeks post-transplantation. Half of the animals in each groups (5 mice per group) were euthanized at 12 weeks post transplantation and bone marrow and terminal blood were collected for analysis. The remaining animals in each group (5 mice per group) were sacrificed at 20 weeks post transplantation.

[0218] Blood collection, cell harvest and processing: Peripheral blood was collected via the submandibular vein or cardiac puncture into EDTA tubes and centrifuged at 500 x g for 5 min to remove the plasma. Following phosphate-buffered saline (PBS) bovine serum albumin (BSA) wash and centrifugation, a 10X volume of hemolytic buffer was added to the pellet, and the mixture was incubated at 37°C for 15 min, centrifuged and washed again. The pelleted fraction was reconstituted in 1 mL PBS BSA; an aliquot was removed and centrifuged at 1,000 x g for 5 min, with the resultant pellet preserved for genotyping. The supernatant fraction was utilized for FACS analyses.

[0219] Bone marrow, femur, tibia and pelvic bones were collected in Iscove's Modified Dulbecco's Medium (IMDM) containing fetal calf serum (FCS); total bone marrow was flushed into a PBS BSA solution and filtered using a 70 µm nylon strainer. Volume was adjusted to 10 mL with PBS BSA, and an aliquot was used for cell counting (Cellometer).

[0220] ZFN activity was analyzed using MiSeq deep sequencing. In brief, Genomic DNA from mice injected with either untransfected control CD34+ HSPC or CD34+ HSPC transfected with enhancer targeting ZFN mRNA was isolated from blood samples obtained at 8 week and 12 week or from bone marrow at 12 weeks post-injection. The region of interest (containing the ZFN binding site within the BCL11A locus) was PCR amplified and the level of modification was determined by paired end deep sequencing on the Illumina platform (MiSeq).

[0221] To generate libraries compatible with the Illumina MiSeq sequencing platform, adaptors, barcodes, and flow cell binder (short DNA sequence) were attached to the target specific amplicons using two sets of fusion primers in sequential PCRs. For MiSeq evaluations of human BCL11A enhancer modification in the mouse blood and bone marrow samples, the protocol had to be adjusted due to the low target DNA amounts in these samples.

[0222] The following primers were used for the MiSeq Adaptor PCR:

PRJIYLFN-f2: ACA CGA CGC TCT TCC GAT CTN NNN AGT CCT CTT CTA CCC CAC CCA (SEQ ID NO:32)
and

PRJIYLFN-r4: GAC GTG TGC TCT TCC GAT CTC TAC TCT TAG ACA TAA CAC ACC AGG G (SEQ ID NO:33).

[0223] For the analysis, DNA from mouse bone marrow samples was isolated by DNeasy and approximately 100 ng of DNA were used in each PCR reaction. DNA from mouse blood samples was isolated by Tissue XS and 10 µL of the 15 µL isolated DNA was used in each reaction. In addition to the DNA, the following were added to each MiSeq PCR reaction: 25 µL HotStar Taq mix (Qiagen), 0.5 µL each of the BCL11A enhancer primers listed above (at a concentration of 100 nM), and water to a 50 µL total reaction volume. Typical MiSeq PCR conditions were: 95°C denaturation for 15', and 30 cycles at 94°C for 30", 62°C for 30" and 72°C for 40", followed by a 10' elongation at 72°C. After the MiSeq PCR, the PCR product was diluted between 1:50 and 1:200 with water, or left undiluted for samples with very low starting cell numbers. Barcode PCR was performed with 1 µL of the MiSeq PCR product diluted as described above, 25 µL HotStar Taq mix, 1 µL forward barcode primer, 1 µL reverse barcode primer (both at a concentration of 10 nM) and water to a 50 µL total reaction volume. Barcode PCR conditions were: 95°C denaturation for 15', and 18 cycles at 94°C for 30", 60°C for 30" and 72°C for 30", followed by a 10' elongation at 72°C. Barcode PCR products were pooled and sequenced on the Illumina MiSeq sequencer. The results are shown in Table 5 above.

[0224] FACS analysis for chimerism, and cell lineage determination. To assess the degree of human chimerism, the fraction of cells in the peripheral blood (at 8, 12, 16 and 20 weeks post engraftment) and bone marrow (at 12 and 20 weeks post engraftment) were stained with hCD45-APC Cy7 (Biolegend) and hCD45-BV510 (BD Biosciences) antibodies respectively and FACS analysis was performed. In addition, hematopoietic lineages analysis was performed by staining bone marrow cells with the specific antibodies described in Table 6 below:

Table 6: Antibody sources for cell markers

CD3-FITC: BD 561807 (clone UCHT1)	BD 561807
CD19-PE: BD 340364 (clone SJ25C1)	BD 340364
CD45-BV510	BD 563204
Lin-APC (CD3/UCHT1, CD14/HCD14, CD16/3G8, CD19/HIB19, CD20/2H7, CD56/HCD56)	BIOLEGEN 348803
CD33-PE-CF594	BD 562492
GlyA-FITC	DAKO 0870
CD38-PerCP Cy5.5	BD 551400
CD14-PE	BD 555398
CD34-PE Cy7	BD 560710
CD71-APC Cy7/H7	BD 563671
CD15-BV650	BD 564232
CD8-PerCP Cy5.5	BD 341051
CD4-PE Cy7	BD 344612
CD56-APC	BD 318310
IgM-APC Cy7	BD 314520
CD20-BV650	BD 563780

[0225] In addition, to purify and sort the HSPC populations, we used an enrichment/depletion strategy using magnetic cell separation (MACS). Bone marrow cells were first stained with CD19-biotin, CD3-biotin, B220-biotin, TER119-biotin and m-ckit-biotin (BD Biosciences) and then incubated with anti-biotin beads (Miltenyi Biotec). The positive fraction and depleted fraction were separated using LS columns (Miltenyi Biotec) placed in the magnetic field of a MACS. After separation, the positive fraction was stained with Streptavidin-APC, CD3-FITC, CD19-PE, CD45-BV510 (BD Biosciences) and the depleted fraction with CD34-FITC (BD Biosciences), Gly-A-PE (DAKO), CD19-APC (BD), Lin-APC (Biolegend), Streptavidin-APC, CD45-BV510, CD33-PE-CF594 (BD) and CD38-PECy-7 (Biolegend).

[0226] Untransfected HSPC and 63014/65722 -transfected HSPC were engrafted into NSG mice using standard procedures as described above. The degree of human chimerism in these mice following engraftment was assessed by measuring the fraction of hCD45 positive cells using FACS.

[0227] Figure 7 shows the percentages of human CD45+ cells in peripheral blood collected at 8, and 12 weeks post-transplant and Figure 8 shows percentages in bone marrow harvested at Week 12. As shown, engraftment levels in this study were comparable human chimerism following engraftment of untransfected control and 63014/65722 transfected HSPC. Only 3 mice out of 60 distributed through the groups did not have CD45+ cells indicating a failure to engraft.

[0228] Reconstitution of various hematopoietic cell lineages was tested by FACS analysis of bone marrow cells obtained at week 12 with antibodies recognizing lineage specific cell surface markers using standard procedures. As shown in Figure 9, comparable representation of all analyzed human hematopoietic lineages in the bone marrow at week 12 post-injection between the BCL11A specific ZFN encoding mRNA treated CD34+ cell progeny and that of the untransfected cells was observed. Bone marrow of the mice sacrificed at Week 12 post-engraftment was isolated and the distribution of various hematopoietic lineages was analyzed by FACS using antibodies recognizing the indicated lineage markers. All numbers are given as the ratio of the cells staining positive for the indicated lineage marker versus the percentage of human CD45 positive cells, except for the cells expressing the erythroid marker Cd71+ (Ter119) in Figure 9C, which are given as the percentage of positively staining cells in the entire population since erythroid cells are not CD45 positive.

[0229] The levels of gene modification at the BCL11A erythroid enhancer (% of alleles with insertions and deletions [indels]) were assessed by deep sequencing of the ZFN target region using the MiSeq sequencing platform as described above. The data are shown in Figure 10 for blood samples from week 8 and week 12, and in Figure 11 for bone marrow samples from week 12 and sorted lineages derived from the week 12 bone marrow cell samples of the 63014/65722 treated cells. For comparison, the indel percentages measured 2 days after the transfection (as listed in Table 5) are also shown on the graphs of Figures 10 and 11.

[0230] In addition, good retention of gene modification at the BCL11A erythroid enhancer was found for both 63014/65722 -treated HSPC donor sets at the various time points and in the various lineages. Comparable modification was observed in both BCL11A dependent (B cells, 'CD19'; primitive progenitors, 'CD38H') and BCL11A independent (myeloid 'CD33') lineages. Although the input gene modification levels were higher in the PB-MR-003 donor sample than in the PB-MR-004 donor sample, the PB-MR-004 derived cells consistently show higher modification levels, *i.e.* better retention of modification, in mice than those derived from PB-MR-003.

[0231] Overall, the observed retention of gene modification at the BCL11A erythroid enhancer in mice was consistent with that observed in prior mouse experiments using a number of ZFNs targeting a variety of gene targets.

[0232] Furthermore, as human erythroid progenitors are not able to differentiate in mice, to determine the

amount of BCL11A targeted gene modification that occurred in these cells, bone marrow cells were removed from the mice and differentiated *in vitro*. In these experiments, bone marrow derived human cells were removed from sacrificed mice at week 12 following engraftment and differentiated *in vitro* as described above. BCL11A target gene modification was measured by high-throughput Miseq sequencing of DNA isolated from cells at day 14 of the differentiation.

[0233] Modification data (indels) are presented in Figure 12, which shows the modification levels at day 14 of the erythroid differentiation. Indel percentages at day 14 of the *in vitro* differentiation vary markedly for each culture that was generated from cells isolated from one mouse, reflecting the oligocellular nature of the expansion obtained under these conditions. The data indicate that BCL11A enhancer modification mediated by the 63014/65722 ZFNs was not markedly altered during the erythroid differentiation. As was observed in the blood and bone marrow samples, erythroid progeny samples of PB-MR-004 derived cells showed higher average levels of modification than erythroid progeny of PB-MR-003 derived cells.

[0234] The relative levels of various globin mRNAs were determined by RT-PCR analysis of RNA isolated from cells at day 14 of the *in vitro* erythroid differentiation, and the data is presented in Figure 13A where the relative γ -globin to β globin mRNA and γ -globin to α -globin mRNA ratios (Figure 13B) averaged out for the 5 erythroid cultures from each group. Both in the untransfected and the 63014/65722 treated samples γ -globin to β globin or γ -globin to α globin mRNA ratios differ widely between the erythroid progenies of individual mice from the same group. The donor PB-MR-004 derived cultures show on average lower γ -globin ratios than those from donor PB-MR-003, in line with the better maturation observed for PB-MR-004 derived samples. However in spite of this variability, the ZFN treated sample averages show an ~ 1.5 -2 fold increase in γ globin mRNA levels compared to their respective untransfected counterparts.

[0235] Globin protein levels were assessed by HPLC analysis. Figure 14 shows globin protein analyses of samples harvested at day 16 of the differentiation. The Gamma globin (sum of the Agamma and Ggamma peaks) to alpha globin ratios were determined, as well as the Gamma globin (sum of the Agamma and Ggamma peaks)/ over beta-like globin ratios (sum of the Agamma, Ggamma, beta and delta-globin peaks) and the averages for each group are shown above each bar. In line with the poor erythroid differentiation of the PB-MR-003 derived samples the gamma-globin levels in the untransfected cells derived from this donor were very high ($\sim 30\%$), and therefore ZFN treatment resulted in only a 1.2 fold increase in gamma-globin levels. The PB-MR-004 showed more typical untransfected levels ($\sim 9\%$) and exhibited an ~ 2 -fold increase in gamma-globin protein levels after 12 weeks passage through the mouse.

[0236] It is thought patients that have a $>8.6\%$ of γ globin naturally are at an advantage as compared to patients with γ globin levels $<8.6\%$ (Platt et al. (1994) N Engl J Med, 330:1639-44). In fact, achieving a chimeric 10-20% percentage of non-sickle cell RBCs through engraftment of edited cells may lead to clinical improvement (Chang et al. (2017) Mol Ther Methods Clin Dev 4:137-148. doi 10.1016/j.omtm.2016.12.009). Thus, despite having to go through an *in vitro* erythroid differentiation process, the percentage of chimeric cells, and the level of γ -globin protein being detected are indicative of therapeutic efficacy.

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Patentkrav

1. Zinkfingernuklease (ZFN), der omfatter aminosyresekvensen ifølge SEQ ID NO:29 og aminosyresekvensen ifølge SEQ ID NO:31.
5
2. Ét eller flere polynukleotider, der koder for én eller flere zinkfingernukleaser ifølge krav 1, eventuelt hvor det ene eller de flere polynukleotider er mRNA.
- 10 3. Ét eller flere polynukleotider ifølge krav 2, der omfatter SEQ ID NO:28 og SEQ ID NO:30.
4. Én eller flere vektorer, der omfatter sekvenser, som koder for én eller flere zinkfingernukleaser ifølge krav 1.
15
5. Celle, der omfatter zinkfingernukleasen ifølge krav 1, det ene eller de flere polynukleotider ifølge krav 2 eller krav 3 eller den ene eller de flere vektorer ifølge krav 4.
- 20 6. Celle ifølge krav 5, hvor:
(i) cellen er en stamcelle eller en forløbercelle, eventuelt hvor cellen er en human celle, og/eller
(ii) cellens genom er modificeret af zinkfingernukleasen, eventuelt hvor den genomiske modifikation er valgt fra gruppen
25 bestående af insertioner, deletioner og kombinationer deraf, såsom en genomisk modifikation inden for +58-regionen af en BCL11A-enhancersekvens.
7. Celle eller cellelinje, der er fremstillet fra cellen
30 ifølge krav 5 eller krav 6, og som omfatter zinkfingernukleasen ifølge krav 1, det ene eller de flere polynukleotider ifølge krav 2 eller krav 3 eller den ene eller de flere vektorer ifølge krav 4.
- 35 8. Delvist eller fuldstændigt differentieret celle, der stammer fra cellen eller cellelinjen ifølge et af kravene 5 til 7, og som omfatter zinkfingernukleasen ifølge krav 1, det ene

eller de flere polynukleotider ifølge krav 2 eller krav 3 eller den ene eller de flere vektorer ifølge krav 4.

9. Celle eller cellelinje ifølge et af kravene 5 til 8, hvor cellens genom er modificeret af zinkfingernukleasen, og cellen eller cellelinjen udviser øget ekspression af gamma- og/eller betaglobin i sammenligning med en celle eller en cellelinje uden den genomiske modifikation.

10. Farmaceutisk sammensætning, der omfatter zinkfingernukleasen ifølge krav 1, det ene eller de flere polynukleotider ifølge krav 2 eller krav 3, den ene eller de flere vektorer ifølge krav 4 og/eller cellen eller cellelinjen ifølge et af kravene 5-9.

11. Farmaceutisk sammensætning ifølge krav 10, der omfatter cellen eller cellelinjen ifølge et hvilket som helst af kravene 5-9 og ydermere omfatter propagerede og differentierede celler, der stammer fra cellen eller cellelinjen.

12. In vitro-fremgangsmåde til modificering af en endogen BCL11A-enhancersekvens i en celle, hvilken fremgangsmåde omfatter indgivelse af zinkfingernukleasen ifølge krav 1 eller det ene eller de flere polynukleotider ifølge krav 2 eller krav 3 til cellen, således at den endogene BCL11A-enhancersekvens modificeres, eventuelt hvorved:

- (i) fremgangsmåden ydermere omfatter indføring af en eksogen sekvens i cellen, således at den eksogene sekvens indsættes i den endogene BCL11A-enhancersekvens, eller
- (ii) modificeringen omfatter en deletion.

13. Farmaceutisk sammensætning ifølge krav 10 eller krav 11 til anvendelse i en fremgangsmåde til behandling af en hæmoglobinopati, såsom en thalassæmi (f.eks. beta-thalassæmi) eller seglcellesygdom hos en patient, hvilken fremgangsmåde omfatter indgivelse af den farmaceutiske sammensætning til patienten, eventuelt hvor patienten er et menneske.

14. In vitro-fremgangsmåde til fremstilling af en genetisk modificeret celle, der omfatter en genomisk modifikation i en endogen BCL11A-enhancersekvens, hvilken fremgangsmåde omfatter
5 de følgende trin:

a) kontakt mellem cellen og det ene eller de flere polynukleotider ifølge krav 2 eller krav 3

b) udsættelse af cellen for betingelser, der bidrager til ekspression af fusionsproteinet fra polynukleotidet, og

10 c) modificering af den endogene BCL11A-enhancersekvens med det eksprimerede fusionsprotein i tilstrækkelig grad til at producere den genetisk modificerede celle
eventuelt hvor fremgangsmåden ydermere omfatter stimulering af cellen med mindst ét cytokin.

15

15. Kit, der omfatter zinkfingernukleasen ifølge krav 1, det ene eller de flere polynukleotider ifølge krav 2 eller krav 3, den ene eller de flere vektorer ifølge krav 4 og/eller cellen eller cellelinjen ifølge et af kravene 5-9.

20

16. Celle eller cellelinje ifølge et hvilket som helst af kravene 5-9 til anvendelse i en fremgangsmåde til behandling eller forebyggelse af en hæmoglobinopati, såsom en thalassæmi (f.eks. beta-thalassæmi) eller seglcellesygdom.

25

17. Celle eller cellelinje til anvendelse ifølge krav 16, hvor cellen er en human stamcelle eller forløbercelle, og cellens genom er blevet modificeret i BCL11A-enhancersekvensen ved hjælp af zinkfingernukleasen, eventuelt hvor:

30 (i) cellerne ekspanderes forud for indgivelse til et individ

(ii) cellerne indgives til individet i en knoglemarvstransplantation, hvor cellerne indpodes, differentierer og modnes in vivo, og/eller

(iii) individet forud for indsættelse af et transplantat, der
35 omfatter cellerne, behandles med en mild myeloablativ procedure eller et kraftigt myeloablativt konditioneringsregime.

DRAWINGS

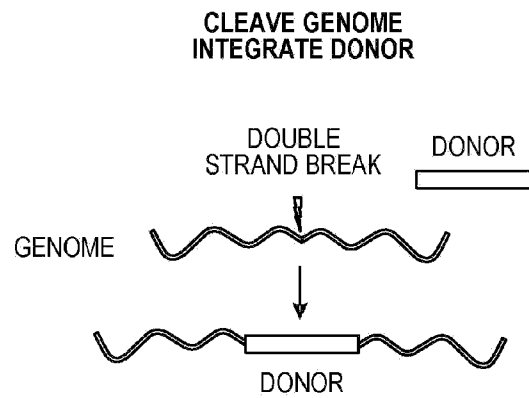


FIG. 1A

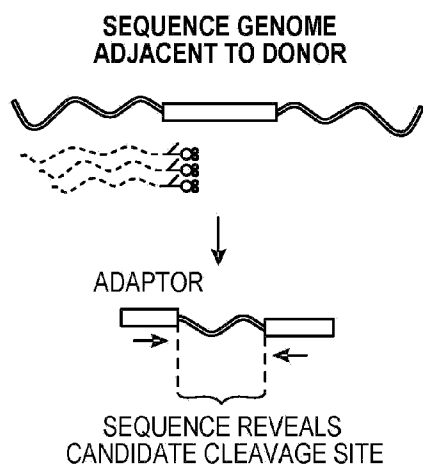


FIG. 1B

CHROMOSOME	BASE #	INTEGRANT COUNT	CHROMOSOME	BASE #	INTEGRANT COUNT	CHROMOSOME	BASE #	INTEGRANT COUNT
2	60495266	1064	22	20508058	15	17	33940450	9
8	119856442	939	1	167637598	14	18	20578334	9
2	62164814	156	1	211655462	14	19	850496	9
10	132654828	105	9	129611194	14	20	29445862	9
14	67422056	102	16	46399310	14	1	51299472	8
4	7829724	93	19	38665086	14	1	201507258	8
2	23702832	88	14	50504	14	2	16715068	8
14	73740920	85	14	50604	14	2	20996616	8
1	29182166	82	2	96873178	13	2	36785314	8
10	69571176	75	2	107763998	13	2	226916064	8
16	15141540	68	2	127820122	13	2	237467876	8
8	94988042	66	6	1596312	13	6	49519624	8
16	2122340	66	6	35468034	13	6	160955686	8
19	45406680	64	6	144064118	13	10	26734304	8
16	67184172	59	8	107386748	13	10	79114636	8
16	46394744	58	10	98910092	13	12	50505884	8
13	75493350	57	12	104048816	13	14	73746866	8
16	34593646	57	14	89466992	13	17	43805774	8
16	46390726	54	1	21635648	12	17	82718744	8
1	54506814	53	2	135453242	12	18	20575274	8
19	55115766	53	6	7899998	12	20	5788510	8
6	53805956	52	8	13105966	12	21	25629778	8
17	29624234	52	9	128604032	12	22	19983370	8
1	26938300	48	11	2992880	12	X	47067946	8
5	145864942	40	13	91576712	12	1	19916860	7
20	37707466	39	14	18295748	12	1	143187676	7
21	33308800	36	16	34587146	12	1	148402426	7
1	204975770	35	21	25607680	12	1	223224616	7
5	132776212	35	22	15226546	12	2	119198990	7
11	33894468	34	1	70765014	11	3	45419400	7
10	101133378	33	1	120942294	11	4	39364694	7
17	28854518	33	1	200702960	11	4	99547294	7
X	30280458	31	3	19942238	11	5	38056212	7
2	84459580	30	5	2057482	11	6	6715482	7
7	131503656	30	6	3196958	11	6	15090744	7
2	112583534	27	9	98755462	11	6	89830096	7
3	49724756	27	11	30406776	11	6	8988012	7
12	55928112	27	1	177943758	10	7	113383212	7

FIG. 2

CHROMOSOME	BASE #	INTEGRANT COUNT	CHROMOSOME	BASE #	INTEGRANT COUNT	CHROMOSOME	BASE #	INTEGRANT COUNT
8	66165666	26	2	218419648	10	8	37700430	7
11	119316888	26	6	136793200	10	8	88289848	7
X	134914730	26	8	25969948	10	9	101660500	7
8	95226212	25	10	46426956	10	9	112857552	7
12	124995298	25	10	47957704	10	10	42097318	7
16	10992070	25	10	52321784	10	10	91591618	7
3	47439806	24	10	52321864	10	10	120431936	7
11	70518294	23	11	102685874	10	11	5225468	7
16	10839466	22	12	54411358	10	12	155086	7
X	66004390	22	12	76936636	10	12	51951306	7
1	9267210	21	15	19782342	10	15	64356576	7
9	91430052	21	16	15220070	10	15	65185434	7
16	30692064	21	16	15220090	10	16	34573028	7
1	155276858	20	16	18131696	10	16	34573058	7
2	173018750	20	16	18131716	10	16	46401528	7
16	88559110	20	18	5447734	10	16	81738062	7
2	149709484	19	19	43779827	10	17	836804	7
8	124924782	19	19	51327816	10	17	75814568	7
13	112237576	19	21	33199996	10	19	17579762	7
20	44683654	19	1	26569750	9	19	38478442	7
1	228965770	18	1	57734650	9	19	40578628	7
7	13991304	18	1	115988958	9	19	41165952	7
7	101110944	18	1	149636460	9	20	56585134	7
11	108224088	18	3	51942092	9	20	36368580	7
12	89874464	18	3	149936952	9	22	36423722	7
1	208159440	17	4	84425848	9	22	2914438	6
5	150115792	17	4	138895946	9	1	91089046	6
12	7096136	16	6	3737388	9	1	115328550	6
15	28018428	16	6	164378486	9	1	125169466	6
19	1279936	16	9	93003938	9	1	125169552	6
20	48636010	16	9	104239026	9	1	155610614	6
1	165379386	15	10	67764450	9	1	155746194	6
3	71103574	15	10	69789200	9	1	201664794	6
5	103961570	15	13	98822320	9	2	131443890	6
7	128077200	15	15	31382604	9	3	4323698	6
8	29495584	15	15	34422898	9	3	58414862	6
16	46399748	15	15	34569124	9	3	66129874	6
16	57534378	15	15	78243618	9	3	87800708	6

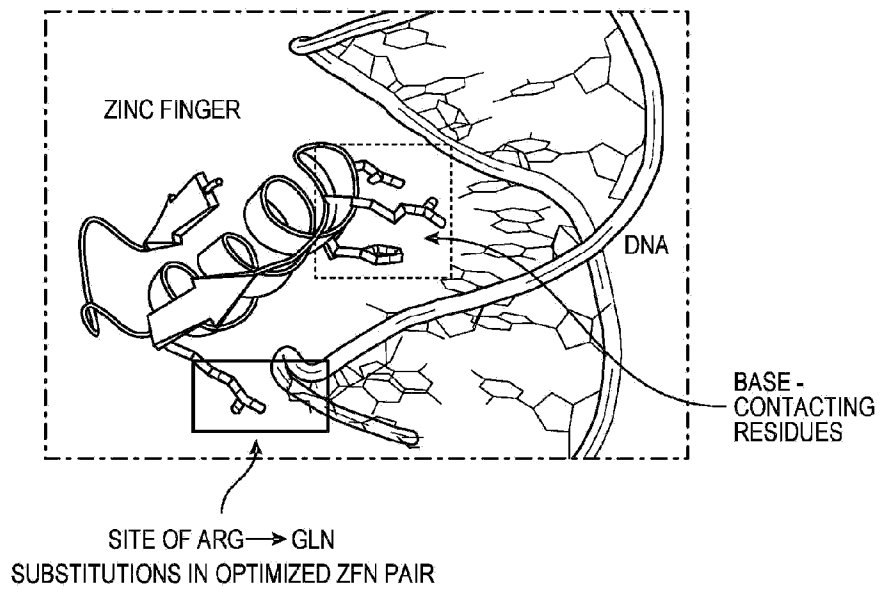
FIG. 2 (cont.)

CHROMOSOME	BASE #	INTEGRANT COUNT	CHROMOSOME	BASE #	INTEGRANT COUNT	CHROMOSOME	BASE #	INTEGRANT COUNT
3	14128736	6	13	79775976	5	9	92277906	4
4	151031930	6	15	40114388	5	9	97934932	4
5	178121148	6	15	68307028	5	9	121566378	4
6	28931180	6	15	82539916	5	9	124392210	4
6	108893514	6	15	86572492	5	9	126591182	4
6	136088666	6	16	21645824	5	9	138212048	4
6	163725808	6	16	53042972	5	10	25357460	4
7	162658	6	16	58873960	5	10	41847448	4
7	35016604	6	16	67841642	5	10	41847504	4
8	30408876	6	16	78808108	5	10	42103514	4
8	91876290	6	17	17025784	5	10	64508254	4
8	119945566	6	17	26884866	5	10	76825618	4
9	4090722	6	17	72902448	5	10	109893420	4
9	104968230	6	18	11456592	5	11	35944902	4
10	72293352	6	18	33538826	5	11	72431102	4
10	79866034	6	19	7159860	5	12	6232920	4
11	118997014	6	19	11922082	5	12	7242432	4
11	121069264	6	19	18462114	5	12	49404046	4
12	3073480	6	19	45686048	5	12	75511410	4
12	26186106	6	20	29881066	5	12	88031364	4
13	43878548	6	20	43558622	5	12	95073610	4
15	41284346	6	20	47575112	5	12	113153372	4
15	58555840	6	22	19545504	5	13	18964478	4
15	60478340	6	22	23284226	5	13	34779902	4
16	3106742	6	14	33684	5	13	113600728	4
16	4518046	6	1	6359676	4	14	92616966	4
16	29292606	6	1	18556502	4	15	33229354	4
16	77989246	6	1	24862574	4	15	64142544	4
16	88266652	6	1	25442520	4	15	67498752	4
16	88732066	6	1	30801948	4	15	75356538	4
17	36938710	6	1	51631166	4	15	76355710	4
19	267174	6	1	72774624	4	15	684294	4
19	39690408	6	1	109213944	4	15	14285910	4
22	40259106	6	1	153609522	4	15	85317250	4
X	40835932	6	1	175878760	4	17	9710280	4
X	1724636	5	1	201154009	4	17	21372082	4
1	6161778	5	1	224687866	4	17	21906776	4
1	17419988	5	2	137774808	4	17	26603948	4

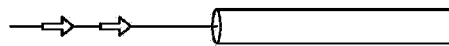
FIG. 2 (cont.)

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1	36401004	5	2	166910124	4	17	26603992	4
1	143208542	5	2	234668446	4	17	40557500	4
1	150528982	5	3	49802584	4	17	43796376	4
1	154750274	5	3	195856500	4	17	49883740	4
2	86563402	5	3	198148214	4	18	13272880	4
2	105038552	5	4	1686620	4	18	30136646	4
2	117954094	5	4	3480800	4	18	30845084	4
2	130230372	5	4	99869180	4	18	74019902	4
2	176188548	5	4	138090616	4	19	40266218	4
3	2041446	5	4	158876704	4	19	42741976	4
3	16078092	5	5	10727038	4	19	42767202	4
3	75669968	5	5	36318878	4	19	42881190	4
3	17898880	5	5	36322678	4	19	49683046	4
3	184689238	5	5	65562414	4	20	4673782	4
4	49119414	5	5	100709408	4	20	24683868	4
4	73573948	5	5	142193948	4	20	44911634	4
4	78700116	5	5	149564992	4	20	48621130	4
5	7057496	5	5	164896712	4	20	49279476	4
5	11767140	5	6	176740968	4	20	49699412	4
5	56314802	5	6	2643624	4	20	56914278	4
5	132314286	5	6	27782516	4	20	62819166	4
6	58116968	5	6	45156450	4	21	10442640	4
6	60870568	5	6	146750040	4	22	22533154	4
7	128792974	5	6	169090710	4	22	22648222	4
7	133081754	5	7	1214740	4	22	22956152	4
9	92835866	5	7	37357350	4	22	23111094	4
9	131373156	5	7	37357420	4	22	23162790	4
10	50701042	5	7	62355868	4	22	24629320	4
10	87382350	5	7	71679460	4	22	32136074	4
11	49652236	5	7	106778616	4	22	45216526	4
11	61612468	5	7	133077880	4	X	6134666	4
11	62795536	5	8	5753270	4	X	39688532	4
11	6727754	5	8	5753290	4	X	52934730	4
11	82032970	5	8	8244554	4	X	71283192	4
11	88966978	5	8	8371950	4	X	90323678	4
12	6326346	5	8	101955394	4	X	108863278	4
12	104138422	5	8	119000446	4	Y	3900438	4
12	121500124	5	9	38068006	4			

FIG. 2 (cont.)

**FIG. 3A**

		SITE OF ARG	GLN			
		SUBSTITUTIONS IN OPTIMIZED ZFN PAIR			BASE-CONTACTING RESIDUES	
LEFT ZFP	F1	VPAAMAERP	FQCRICMNF	S	DQSNLRHIRTH	
	F2	TGEKP	FACDICGRKF	A	RNFS:LTMTKIH	
	F3	TGSQKP	FQCRICMNF	S	STGN:LTNHIRTH	
	F4	TGEKP	FACDICGRKF	A	TSGS:LTRHTKIH	
	F5	THPRAPIPKP	FQCRICMNF	S	DQSNLRHIRTH	
	F6	TGEKP	FACDICGRKF	A	AQCC:LFHTKIH	LRGS
RIGHT ZFP	F1	VPAAMAERP	FQCRICMNF	A	RNDHRTHTHTKIH	
	F2	TGEKP	FQCRICMNF	S	QKAHLIRHIRTH	
	F3	TGEKP	FACDICGRKF	A	QKGT:LGEHTKIH	
	F4	TGSQKP	FQCRICMNF	S	RGRD:LSRHIRTH	
	F5	TGEKP	FACDICGRKF	A	RRDN:LHSHTKIH	LRGS

**FIG. 3B**

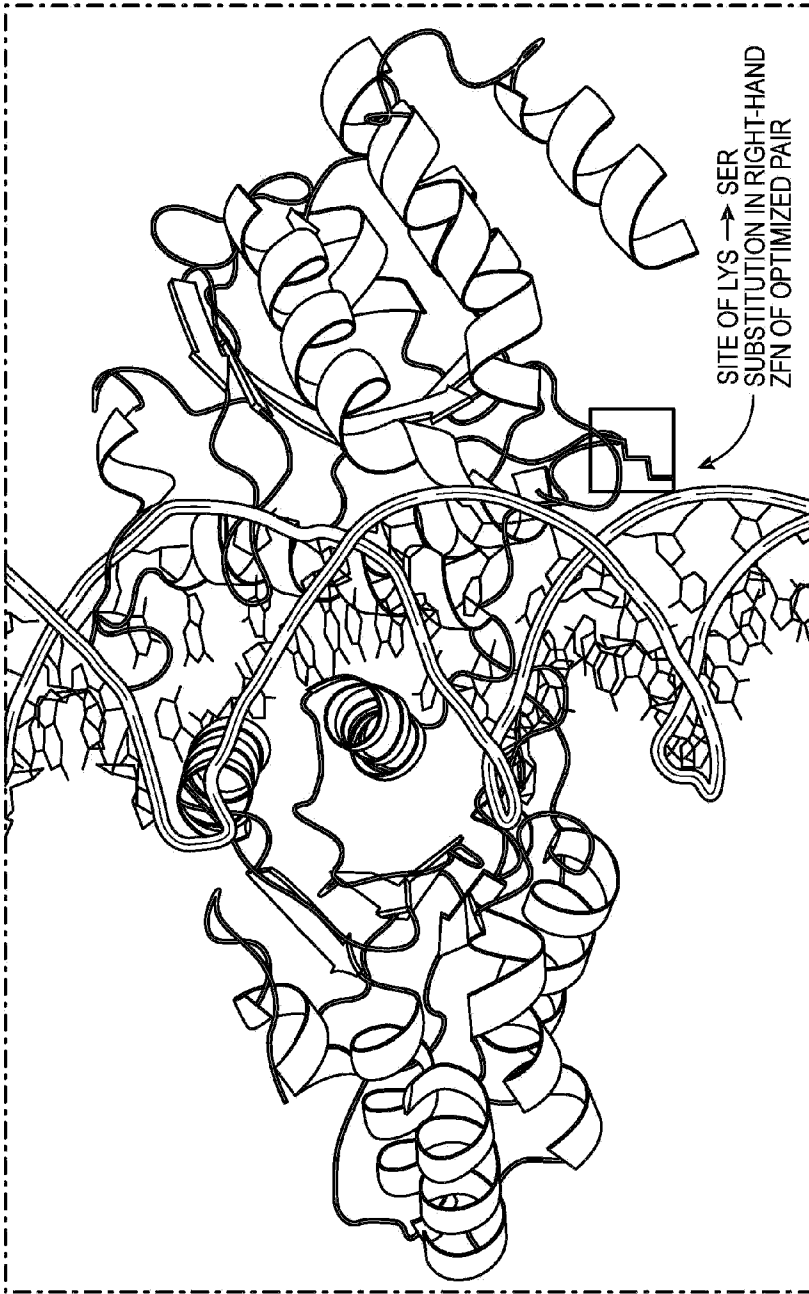


FIG. 3C

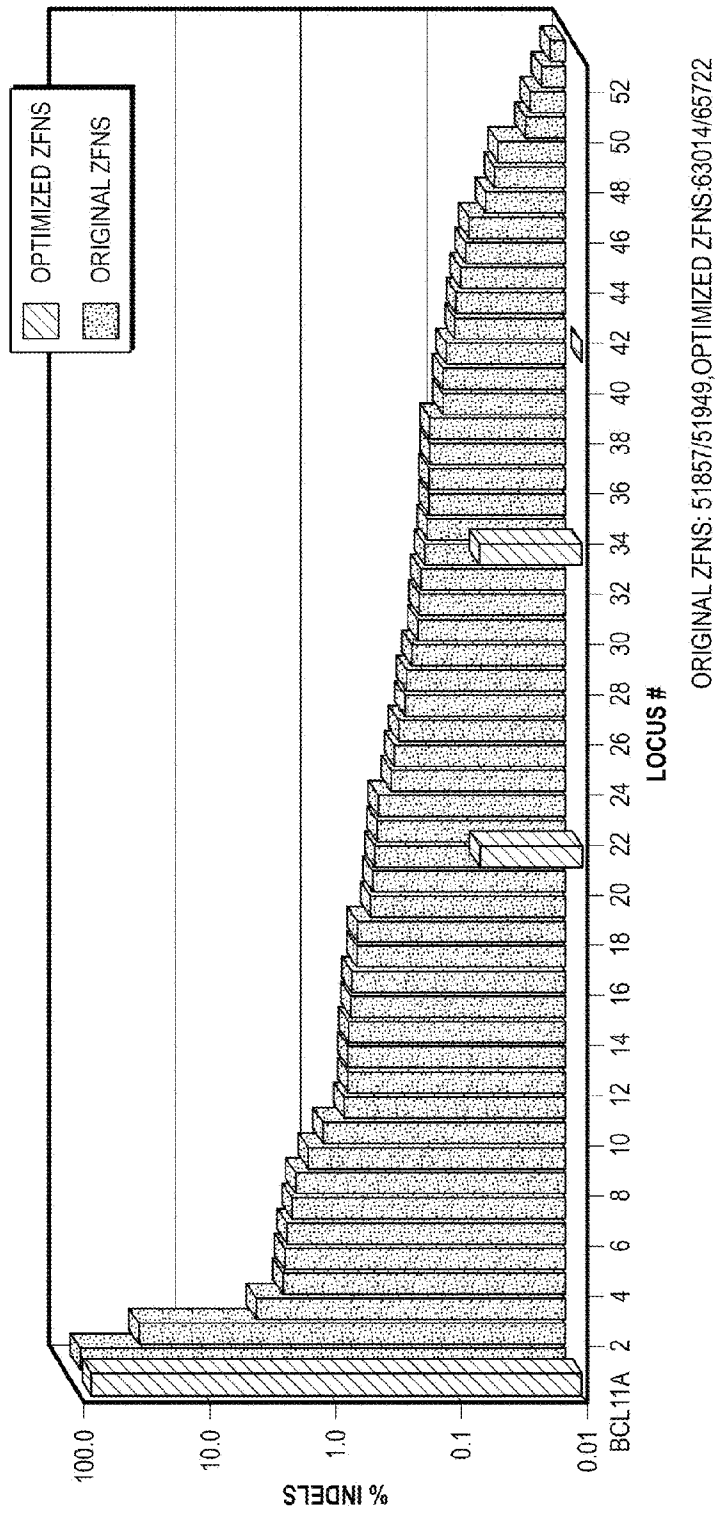
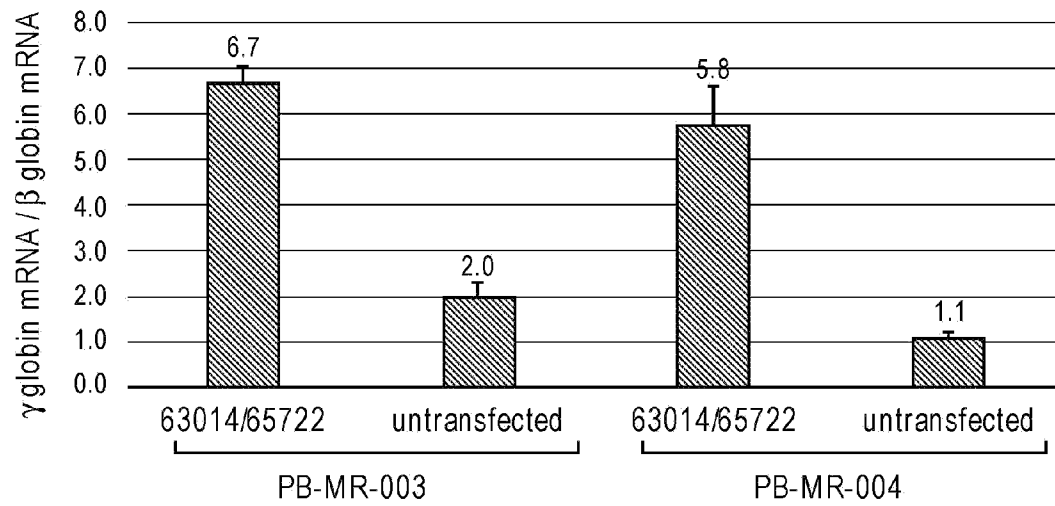
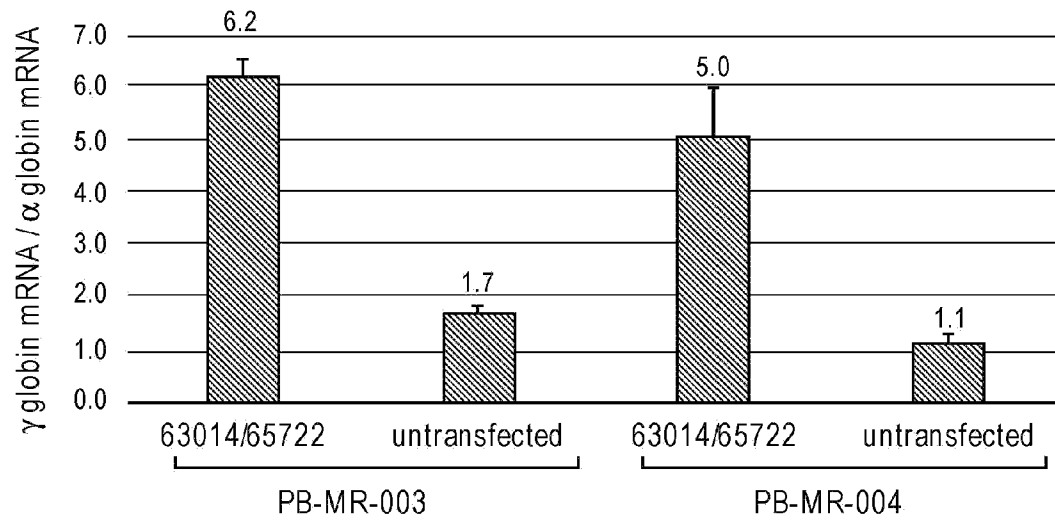
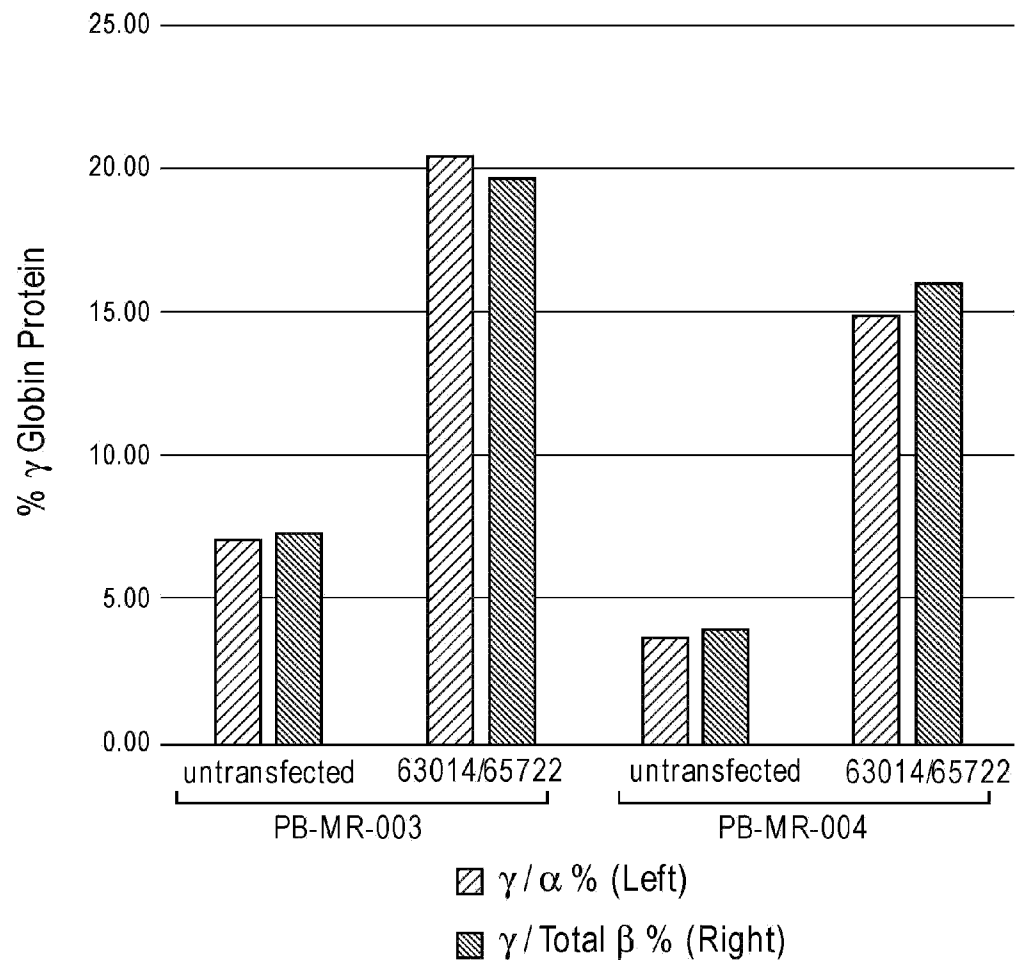
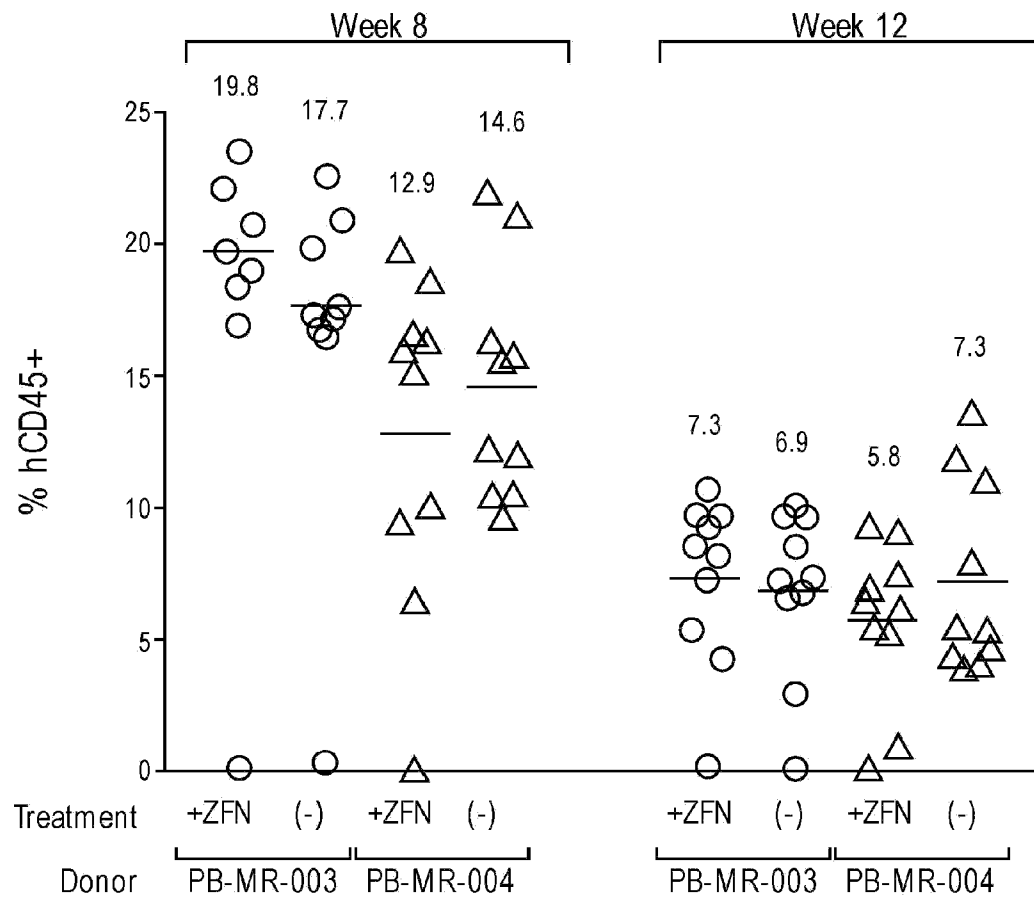
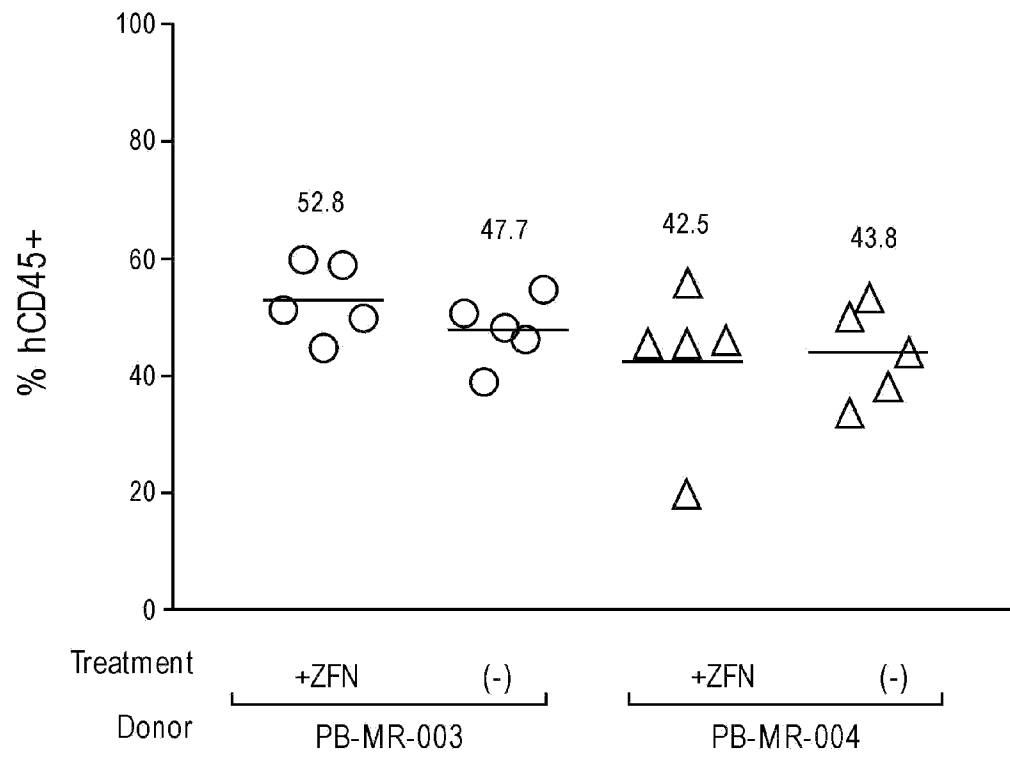


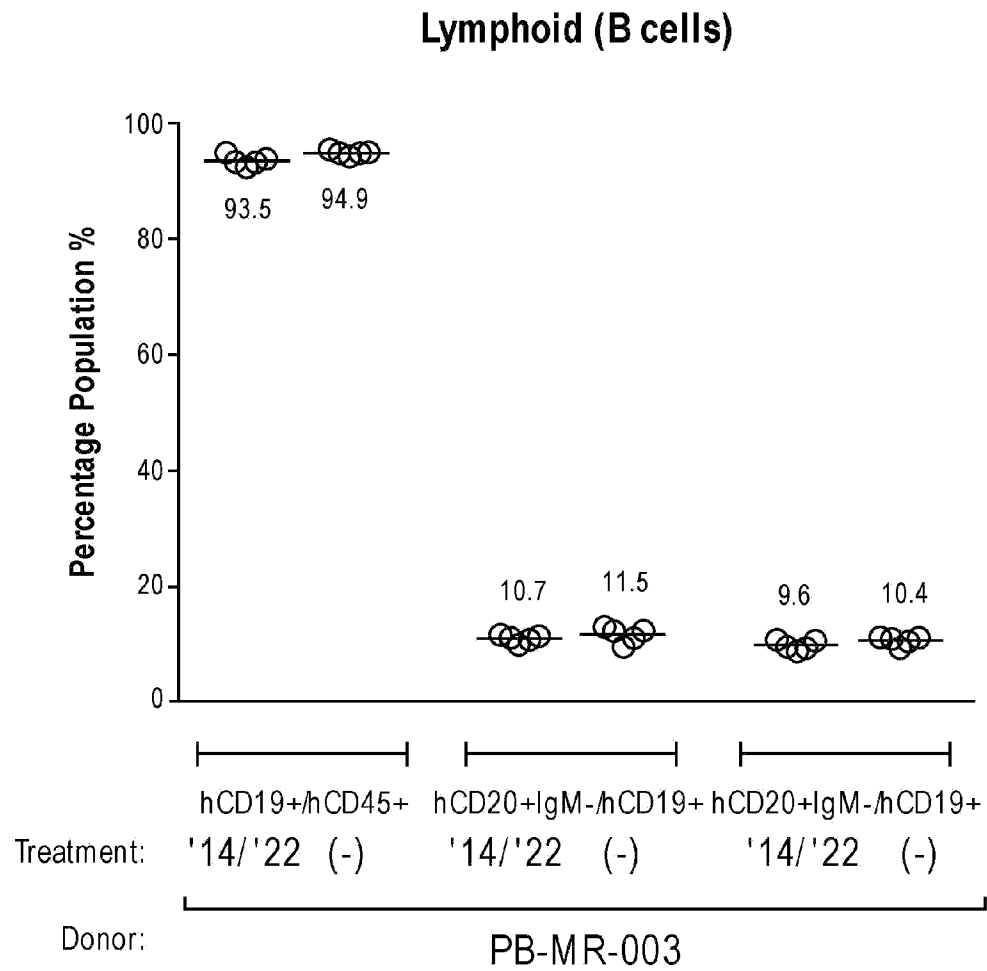
FIG. 4

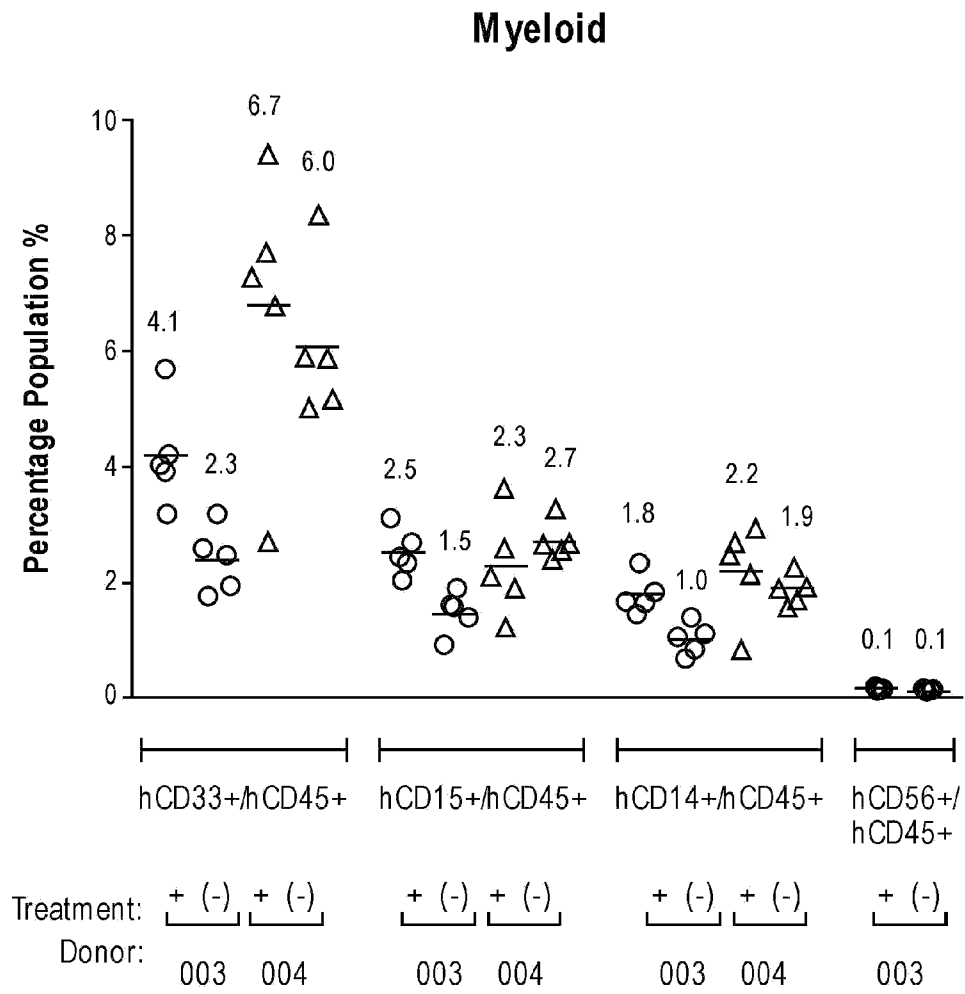
**FIG. 5A****FIG. 5B**

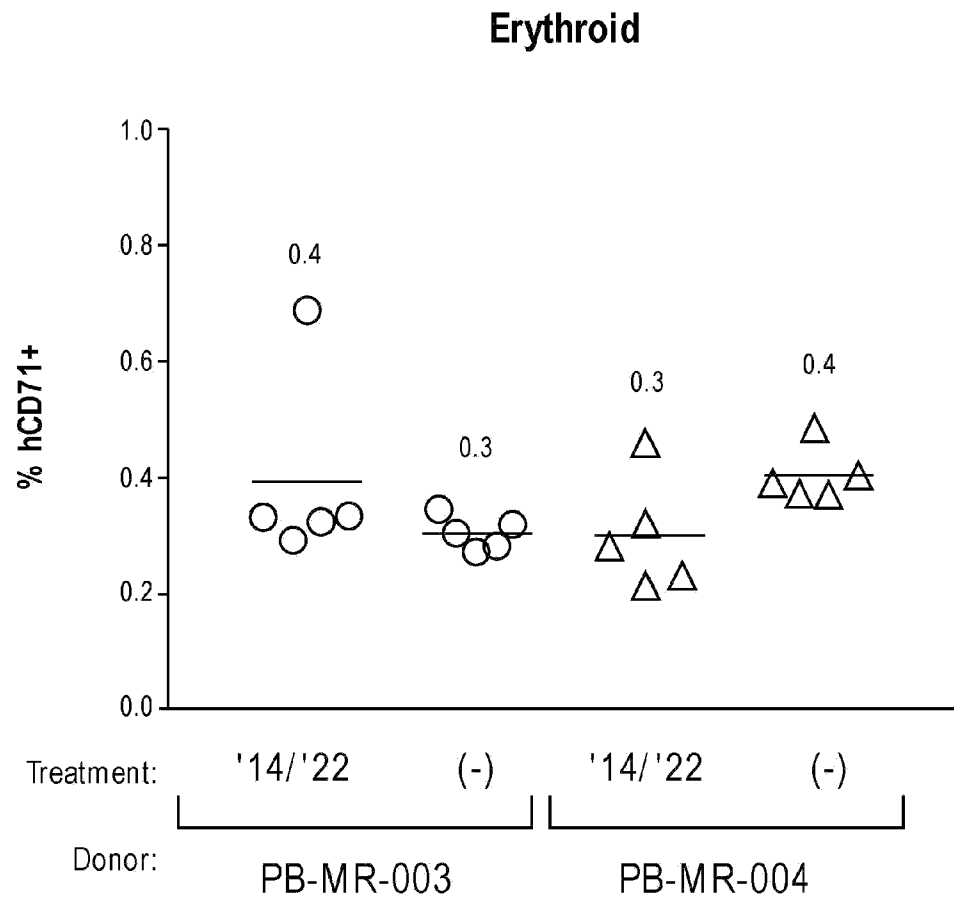
**FIG. 6**

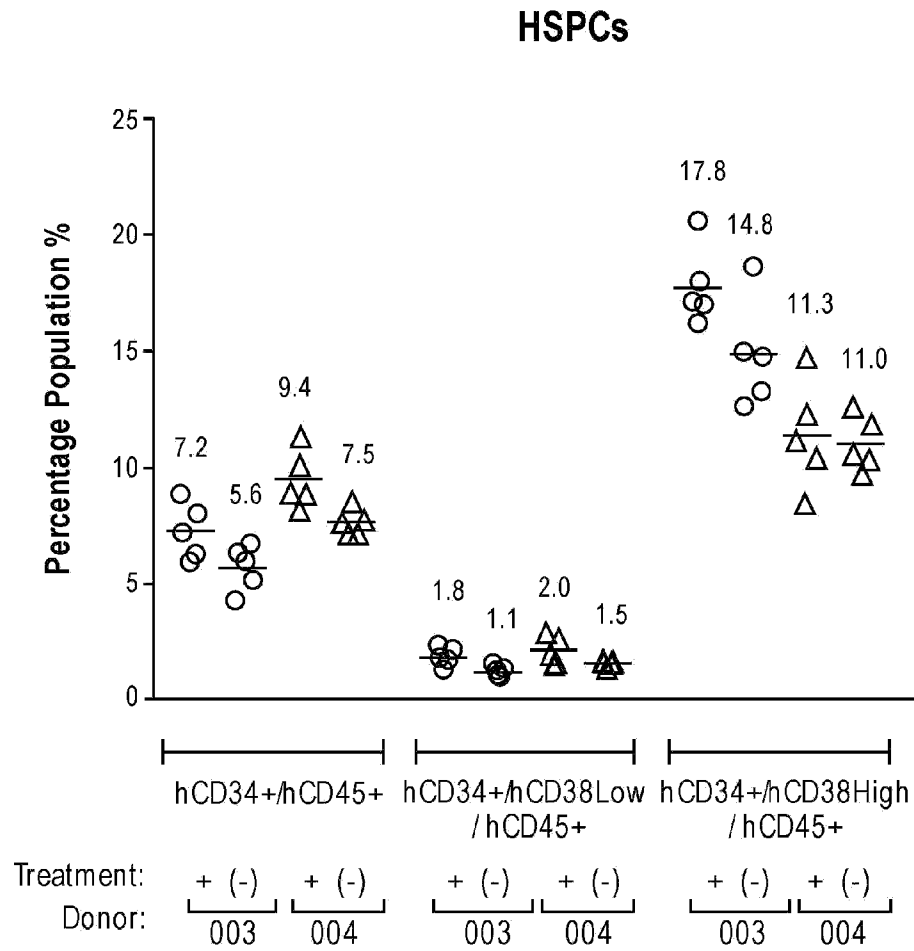
**FIG. 7**

**FIG. 8**

**FIG. 9A**

**FIG. 9B**

**FIG. 9C**

**FIG. 9D**

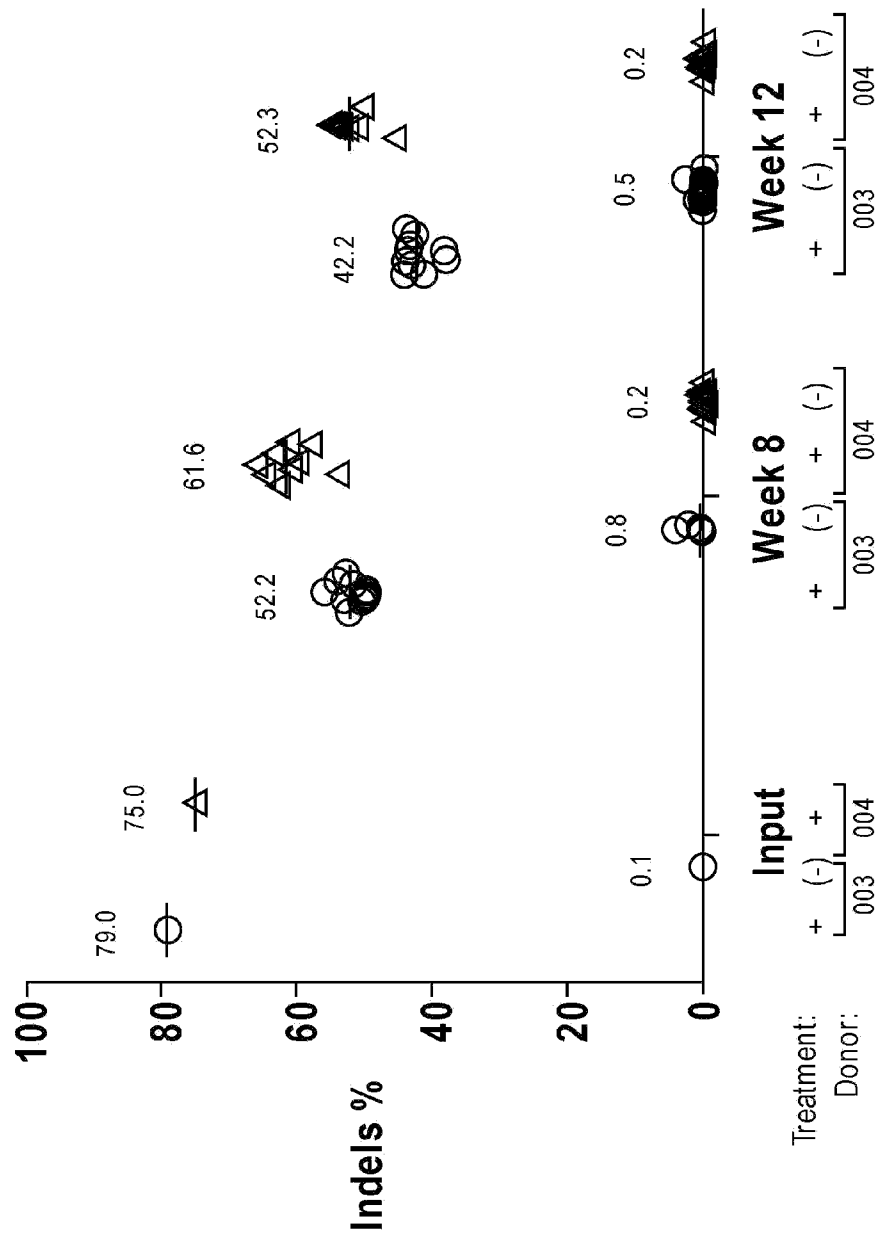


FIG. 10

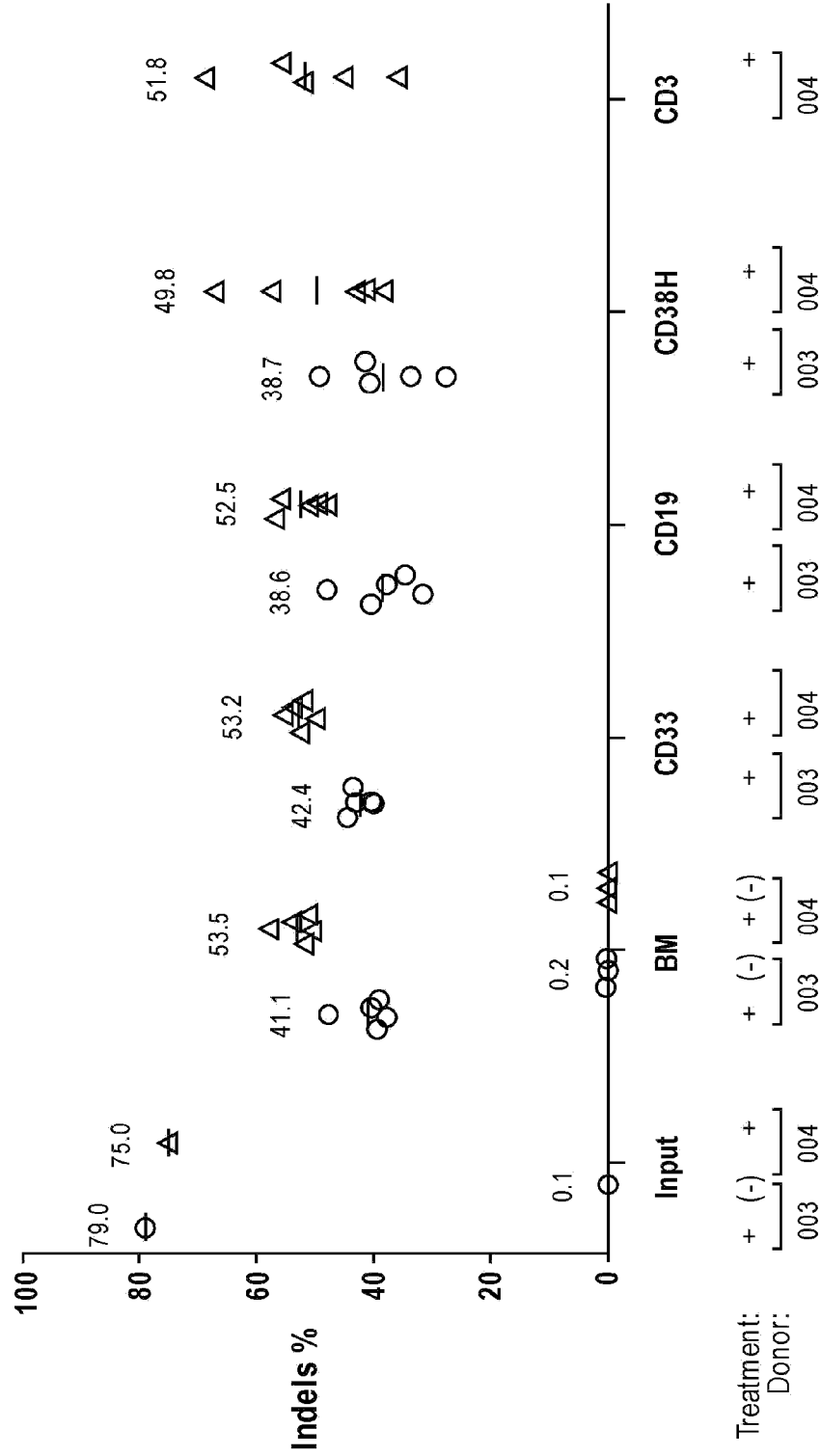
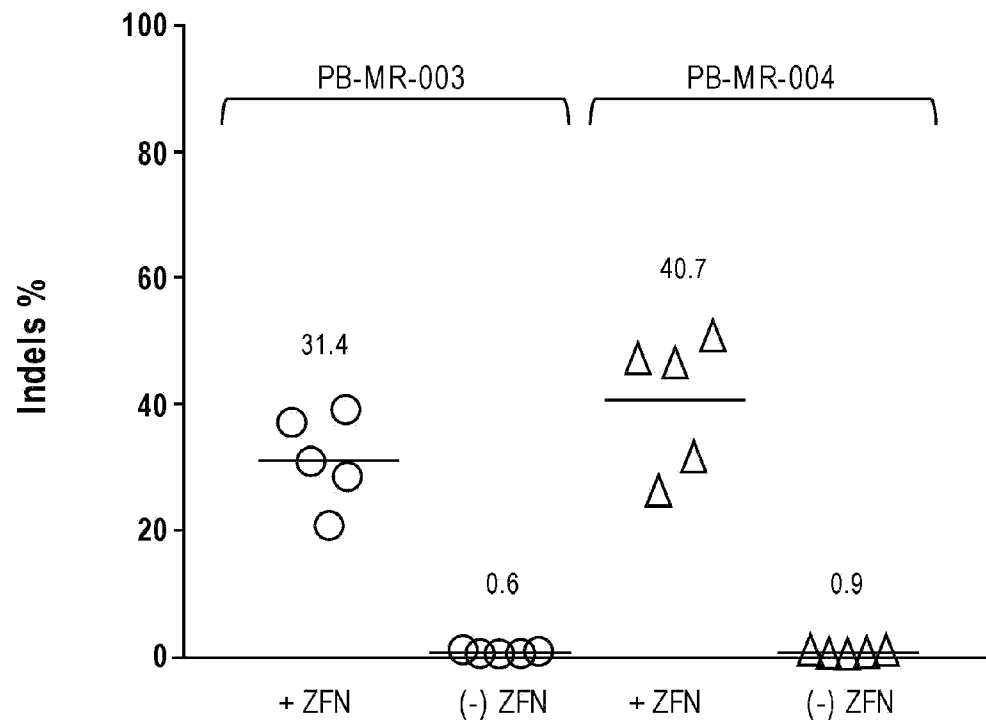
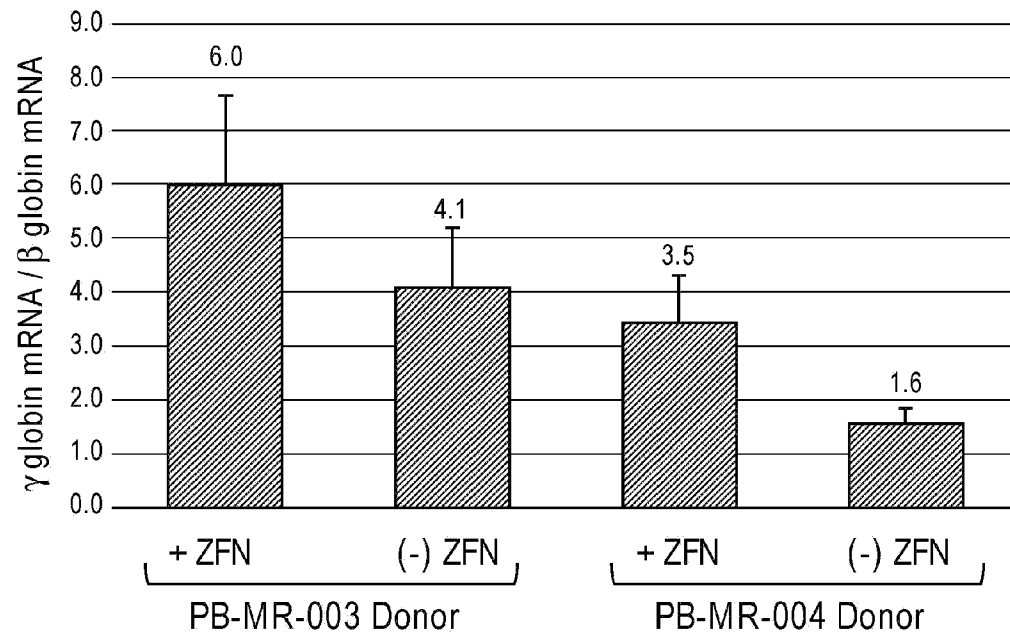
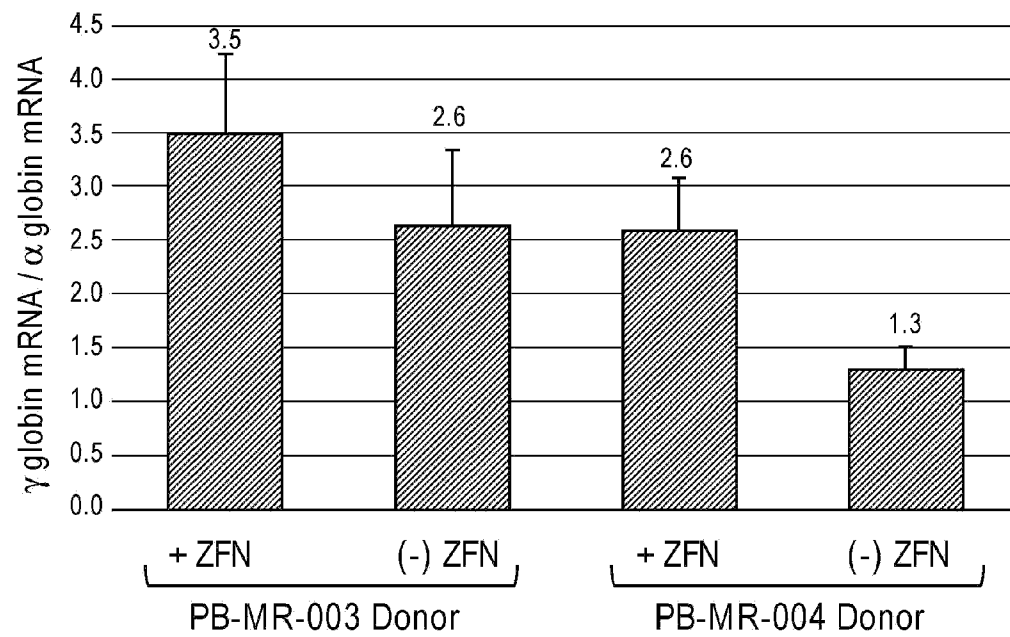
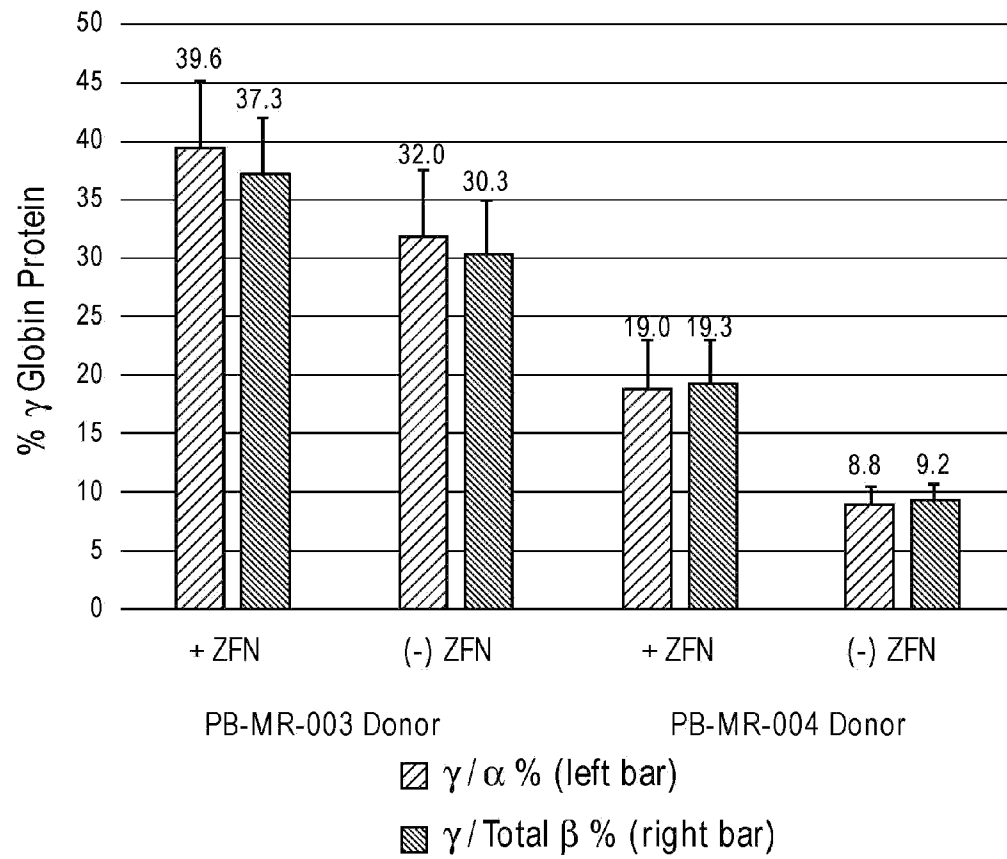


FIG. 11

**FIG. 12**

**FIG. 13A****FIG. 13B**

**FIG. 14**