βT87Q-Globin Gene Therapy Reduces Sickle Hemoglobin Production, Allowing for Ex Vivo Anti-sickling Activity in Human Erythroid Cells

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Lentiviral addition of βT87Q-globin, a modified β-globin with an anti-sickling mutation, is currently being used in gene therapy trials for sickle cell disease (SCD) and β-thalassemia patients. βT87Q-globin interferes with sickle hemoglobin (HbS) polymerization. Here, we generated the SCD mutation in an immortalized human erythroid cell line (HUDEP-2) to investigate the anti-sickling activity of βT87Q-globin. Sickle HUDEP-2 (sHUDEP-2) cells produced robust HbS after differentiation and sickled under deoxygenated conditions, comparable with SCD CD34+ progeny. Lentiviral transduction provided 9.5–26.8 pg/cell βT87Q-globin (R² = 0.83) in a vector copy number (VCN)-dependent manner, resulting in a significant reduction of sickling ratios (R² = 0.92). Interestingly, βT87Q-globin transduction markedly reduced endogenous βγ-globin (R² = 0.84) to an undetectable level (0.4–16.8 pg/cell) in sHUDEP-2 cells, as well as endogenous β-globin in human CD34+ cell-derived erythroid cells. RNA sequencing (RNA-seq) analysis with βT87Q-transduced sHUDEP-2 and human CD34+ cell-derived cells revealed activation of inflammation- and proliferation-related programs, suggesting minimal changes in background gene expression except for βT87Q-globin expression and endogenous β/βγ-globin suppression. In summary, using sHUDEP-2 and CD34+ -derived cells, we demonstrated that lentiviral addition of βT87Q-globin strongly reduced endogenous β-/βγ-globin expression, resulting in an anti-sickling effect. Our findings should be helpful to understand the anti-sickling effects of therapeutic genes in SCD gene therapy.

INTRODUCTION

Sickle cell disease (SCD) is a monogenic red blood cell (RBC) disorder with approximately 300,000 new affected individuals globally per year, which is predicted to rise to 400,000 patients in 2050.1 A homozygous missense mutation (E6V, rs334) in the β-globin gene leads to polymerization of hemoglobin (Hb) under deoxygenated conditions that results in various secondary molecular and cellular changes, many of which disrupt circulation and contribute to major tissue damage.2 As of now, allogeneic hematopoietic stem cell (HSC) transplantation is the only curative option available to patients outside of clinical trials. Despite the procedure’s high success rate, a majority of sickle cell patients lack a histocompatible sibling donor, and there remains a significant risk for graft-versus-host disease (GVHD), graft rejection, and transplant-related mortality.3 Thus, the development of novel and efficient treatment modalities is of great interest to the biomedical community.

In SCD, sickle Hb (HbS) forms polymers under deoxygenated conditions that drastically change mechanical and rheological properties of RBCs, interrupting circulation and reducing oxygen transfer to vital organs. Efforts to develop gene therapy for SCD include various anti-sickling gene transfer approaches (including β-globin, γ-globin, and β/γ-globin hybrids) in SCD blood progenitor cells (reviewed in Demirci et al.4). However, insufficient globin expression was observed in early attempts because of ineffective transfer of large cassettes, instability of globin expression from cassettes in retroviral vectors, and a lack of long-term persistence of erythroid-specific globin expression.5,6

The inhibitory effect of β-globin on sickling is relatively low compared with γ-globin.6,9 Biochemical studies revealed that a critical difference at position 87 (Q replacing T) in γ-globin is one of the main reasons for its superior anti-sickling activity.10 Lentiviral delivery of βT87Q-globin containing this anti-sickling mutation has been tested in SCD mouse models; along with an erythroid-specific accumulation of anti-sickling protein, inhibition of sickling and correction of hematological parameters in mice were reported.11 βT87Q-globin transfer is now being evaluated in clinical trials for β-thalassemia and SCD12 (ClinicalTrials.gov: NCT03207009, NCT02906202, NCT01745120, and NCT02151526), and encouraging results have been reported. However, vector copy numbers (VCNs) and transgenic protein amounts are variable, and
A consensus on the protein level needed for therapeutic effects has not yet been established.

Rapid and remarkable advances in molecular biology, particularly genome-editing tools, have led to efficient editing of \(g\)-globin regulatory elements\(^ {13,14}\) and even correction of the SCD mutation\(^ {15}\) in HSCs, generating excitement about the emergence of new therapeutic modalities. However, these approaches should be investigated, optimized, and verified in \(ex\ vivo\) SCD models regarding their molecular mechanism, efficacy, and safety before testing in animal models and subsequent clinical trials to enhance the success rate for early participants in those trials. In terms of speed and cost, \(ex\ vivo\) cell culture models have a great advantage over human primary cells for drug candidate screening and molecular mechanism evaluation. To the best of our knowledge, there is no publicly available SCD cell line for \(ex\ vivo\) research. Here, we introduced the SCD mutation into a previously generated immortalized erythroid progenitor cell line (HUDEP-2)\(^ {16}\) using the CRISPR-Cas9 approach, allowing us to evaluate the anti-sickling activity of \(\beta\)T87Q-globin, as well as its potential mechanism of action using RNA sequencing (RNA-seq) in this cell line.

**RESULTS**

**Sickle HUDEP-2 (sHUDEP-2) Cells Produce Sickle Hemoglobin (HbS)**

(A) Hemoglobin (Hb) electrophoresis of differentiated cells derived from wild-type HUDEP-2 and cells electroporated with ribonucleoprotein complex and donor template containing the sickle cell disease (SCD) mutation (Edited).

(B) \(qRT-PCR\) analysis of single-cell cloned electroporated cells.

(C) Hb electrophoresis for single-cell cloned sHUDEP-2 cells.

(D and E) Cell number (D) and cell surface marker (GPA, CD71, and CD36) expression change (E) during red blood cell (RBC) differentiation of sHUDEP-2 cells (n = 3).

(F) Giemsa-wright staining of sHUDEP-2 cells at day 10 of differentiation.

sHUDEP-2 cell numbers increased over the course of a 14-day differentiation period (Figure 1D), similar to the parental HUDEP-2 cell line as reported previously.\(^ {16}\) There was a \(\sim\)13-fold increase in cell number at day 10 and a \(\sim\)24-fold increase at day 14 of differentiation. sHUDEP-2 cells were evaluated for erythrocyte marker (CD36, CD71, and glycophorin A [GPA]) expressions throughout differentiation. Most of the cells were already positive for CD36 (82.3% \(\pm\) 2.8%), CD71 (68.0% \(\pm\) 2.8%), and GPA (69.0% \(\pm\) 2.9%) at day 0 (Figure 1E), similar to wild-type HUDEP-2 cells.\(^ {16}\) Although there was a slight reduction in CD71 and CD36 expression during the early phase of differentiation, expression increased after 5 days. Moreover, GPA levels reached 99.2% \(\pm\) 2.9% at day 10 of differentiation. Because GPA is a terminal marker for erythrocyte differentiation, we used 10 days for the differentiation experiments. Whereas GPA positivity was almost 100% at day 10 of differentiation, enucleation efficiency was only 6.5% \(\pm\) 2.9% as determined by flow cytometric analysis of cells stained with Hoechst 23322 dye (Figure 1F), which is similar to the enucleation efficiency of wild-type HUDEP-2 cells.
These observations indicate that sHUDEP-2 cells are similar to wild-type HUDEP-2 cells except for the SCD mutation.

**βT87Q-Globin Addition Reduces βS-Globin Production and Sickling in sHUDEP-2 Cells**

Next, we investigated whether sHUDEP-2 would sickle under deoxygenated conditions. Concurrently, we introduced βT87Q-globin gene into the sHUDEP-2 cells using lentiviral transduction at increasing multiplicities of infection (MOIs; 0.5, 1, 2, 5, 10, and 25) to evaluate its anti-sickling activity. Transduced cells were single-cell cloned to obtain a homogeneous VCN inside the cell population. Single-cell-derived clones were subjected to qPCR and 10 clones (with VCNs of 0.6, 1.12, 1.57, 2.45, 3.48, 4.46, 5.25, 6.02, 7.99, and 10.06) were selected for further evaluation. Hb electrophoresis and reverse-phase high-performance liquid chromatography (RP-HPLC) analysis demonstrated significant expression of βT87Q-globin in transduced sHUDEP-2 cells (Figures 2A and 2B). Up to a VCN of 3.48, the βT87Q-globin amount was 0%–36.7%, whereas the cells with a VCN of 3.48 or greater expressed 81.3%–89.2% βT87Q-globin and 1.16%–11.9% HbS. Absolute quantification of Hb in each clone showed that there is a significant polynomial correlation between VCN and βS-globin (R² = 0.8423) and βT87Q-globin (R² = 0.8338) protein levels (Figure 2C). It is also noteworthy that higher βT87Q-globin levels reduced the amount of βS-globin expression in sHUDEP-2 cells at the protein level, indicating an internal feedback mechanism for globin chain incorporation for this cell line or more rapid βT87Q-globin production from the vector construct than βS-globin.

βT87Q-globin-expressing cells were evaluated for their sickling potential under deoxygenated conditions. RBCs derived from patients with a genotype of HbAS (AS) globin rarely sickled, whereas a significant proportion of RBCs with HbSS (SS) genotype sickled under low oxygenic conditions (Figure 2D). Interestingly, a large majority of deoxygenated sHUDEP-2 cells also sickled (51%, 47%, and 42% in 410, 354, and 298 mOsm conditions, respectively). There were clear signs of sickle fiber formation and deformation of the cell membrane, but sickling was irregular and distinct from that observed in patient-derived RBCs. However, the sickling fraction was significantly negatively correlated with VCN (R² = 0.61–0.73) and βT87Q-globin protein amount (R² = 0.92) (Figures 2E, 2F, and S1). Osmolarity of saline solutions strongly affects both the rate and the degree of deformation of deoxygenated sickle cells. Increased osmolarity leads to dehydration of RBCs and increased intracellular HbS concentration, thereby increasing the rate of polymerization under deoxygenated conditions. In this report, we evaluated the influence of VCN on the sickling of HUDEP-2 cells at different osmolarities, as can be seen in Figure 2E. The sickling ratio was 12%–17% for all osmolarity conditions in sHUDEP-2 with a VCN of 10 (Figure 2E), confirming the inhibitory effect of βT87Q-globin on sickling in sHUDEP-2 cells.

To evaluate globin chain RNA and protein levels in both sHUDEP-2 and human CD34⁺-derived cells, we first transduced the cells with a βT87Q-globin vector at different MOIs (Figure 3A). After differentiating transduced cells, RNA and protein levels (normalized to x-globin level) revealed that whereas the transcript copies for
βT87Q-globin were as high as 3.9 ± 0.002 copies per β-globin transcript, the β-globin transcript was as low as 0.25 ± 0.003 copies in sHUDEP-2 cells (Figure 3B). In addition, α-globin RNA levels were similar in βT87Q-globin vector-transduced cells and non-transduced mock control (Figure S2). In line with RNA levels, higher MOI correlated with enhanced βT87Q-globin protein levels (44.9%–87.5%) and reduced amount of HbS (7.3%–47.0%) in sHUDEP-2 cells (Figures 3C and 3D). Similar to sHUDEP-2 cells, healthy donor CD34+-derived cells transduced with βT87Q-globin-expressing lentivirus expressed fewer endogenous β-globin transcripts per α-globin transcript (0.52–0.86 copies per α-globin), while the non-transduced control expressed 1.02 ± 0.03 copies per α-globin (Figure 3E). Accordingly, transduced CD34+-derived cells expressed enhanced βT87Q-globin protein levels (13.3%–34.5%) and lower levels of endogenous β-globin (62.3%–83.0%), supporting the notion that βT87Q-globin expression affects endogenous β- or α-globin expression both at the RNA and the protein levels.

**Distinct Gene Expression Signatures in sHUDEP-2 Cells versus CD34+-Derived Cells Expressing βT87Q-Globin**

We performed gene expression profiling by RNA-seq to determine which transcripts were affected by overexpression of βT87Q-globin. sHUDEP-2 cells and healthy donor mobilized CD34+-derived cells were transduced with control lentiviral (GFP) vector or βT87Q-globin vector. CD34+ cells were transduced at day 2 of our two-phase erythroid culture system, and sHUDEP-2 cells were transduced in expansion medium followed by erythocyte differentiation as described above. At day 7 post-transduction (day 9 of culture for CD34+ cells and day 6 of culture for sHUDEP-2 cells), RNA was isolated for RNA-seq and differential gene expression profiling, and expression data were analyzed by ingenuity pathway analysis (IPA). To confirm the RNA-seq data, we analyzed expression of selected genes in sHUDEP-2 cells and mobilized PB CD34+-derived cells by qRT-PCR. Our results show that Smoothened, Frizzled Class Receptor (SMO), and Adenomatous Polyposis Coli Protein 2 (APC2) are upregulated in CD34+ cells transduced with βT87Q-globin lentiviral vector (Figure S3B).

Using a 1.4-fold change in expression as a cutoff, RNA-seq identified 748 differentially expressed transcripts in sHUDEP-2 cells transduced with GFP expression control vector (VCN = 30.3) and 761 transcripts in sHUDEP-2 cells transduced with βT87Q-globin lentiviral vector (VCN = 11.2) compared with non-transduced (Mock) sHUDEP2 cells. Compared with non-transduced control CD34+ cells, 711 transcripts were differentially expressed in CD34+ cells transduced with GFP control vector (VCN = 1.4) and 725 transcripts in CD34+ cells transduced with βT87Q-globin lentiviral vector (VCN = 0.5), respectively (Tables S1A–S1D). Of these transcripts, 207 were common to both sHUDEP-2 transduced with GFP control vector or βT87Q-globin lentiviral vector (Table S1E), and 307 were common to CD34+ cells transduced with control vector or βT87Q-globin lentiviral vector (Table S1F). The differentially expressed transcripts common to sHUDEP-2 cells transduced with GFP vector or βT87Q-globin vector, or the transcripts common to CD34+ cells transduced with GFP vector or βT87Q-globin vector are most likely non-specific, i.e., an effect of the viral transduction. Interestingly, only seven genes were found to be common to all four groups, including the phospholipase D family member 4 (PLD4) gene, which is involved in inflammatory cytokine responses and regulation of...
nucleic acid sensing (Table S1G). This small overlap in commonly expressed genes among all four groups likely represents the biological difference between the sHUDEP-2 cell line and primary CD34+ cells. In addition, principal-component analysis (PCA) revealed that CD34+ and sHUDEP-2 cells cluster differently (Figure S3C), indicating the different transcriptomic structures of the cells.

Further analysis of the datasets showed that 554 transcripts were specific to sHUDEP-2 cells transduced with βT87Q-globin vector when compared with sHUDEP-2 transduced with GFP control vector (Table S1H), and 418 transcripts were specific to CD34+ cells transduced with T87Q-globin lentiviral vector (Table S1I) when compared with CD34+ cells transduced with GFP control vector. Our IPA of sHUDEP-2 cells showed a predominant cancer-related gene expression signature in both GFP control and βT87Q-globin lentiviral vector groups in comparison with mobilized CD34+-derived cells (Figure 4A). These signatures may reflect the status of the sHUDEP-2 cells because they are immortalized with an inducible HPV16 E6/E7 viral vector system. Analysis of genes specifically expressed in sHUDEP-2 cells transduced with βT87Q-globin vector showed enrichment of genes and pathways involved in amino acid metabolism and protein synthesis, whereas mobilized CD34+-derived cells transduced with βT87Q-globin vector showed preferential expression of genes involved in Wnt signaling (Figure 4B). Wnt signaling is critical in development and stem cell function but has not been shown to have a clear role in globin expression.

Further, the IPA shows that amino acid metabolism, cancer, and small-molecule biochemistry are the most significantly affected network in sHUDEP-2 cells transduced with the βT87Q-globin vector (Figure 5). According to the Bloodspot database, MAPK11, TRB3, and PSAT1 are expressed in erythroid cells, but the majority of the genes in the network do not appear to be normally significantly expressed in that lineage. In mobilized CD34+ cells transduced with the βT87Q-globin vector...
vector, the most significantly affected network is organismal injury and abnormalities, behavior, and gastrointestinal disease (Figure 4B). A significant fraction of the upregulated genes in the network is comprised of small nuclear ribonucleoproteins (snRNPs), which are involved in RNA splicing, i.e., removal of introns from pre-mRNA. Taken together, transduction of sHUDEP-2 cells and mobilized CD34+ cells with βT87Q-globin vector activates common gene expression programs (inflammation) and cancer-related (sHUDEP-2 cells) or developmental/stem cell-related (CD34+ cells) programs.

DISCUSSION

Immortalized cell lines are critical ex vivo models in the study of cellular responses to specific stimuli and basic cellular mechanisms, including differentiation and disease development, because they are cost-effective, easy to handle, unlimited in supply, not associated with ethical concerns, and provide consistent starting materials. Only a handful of cell lines are able to express adult β-globin and are available for RBC metabolism and disorder research, including the HUDEP-216 and BEL-A21 cell lines. In the present study, we introduced the sickle mutation into the widely studied HUDEP-2 cells using CRISPR-Cas9 technology to generate an HbS-expressing cell line. Cell proliferation is thought to enhance homology-directed repair (HDR) in genome editing, as well as HbF induction in erythroid progenitors. The HUDEP-2 cell line contains HbF-expressing subclones, which were selected using lower-concentration stem cell factor (SCF) when thawed from a frozen stock (data not shown). The SCF-oversensitive HbF-expressing clones seem to be more proliferative; thus, biallelic HDR might be more frequently observed in these HbF-expressing clones because of their increased proliferation (Figure 2B). The sickle cell line can be used not only for understanding erythropoiesis, but also to perform SCD mutation correction, HbF induction, and gene addition research. Here, we have shown proof of concept of their ex vivo research potential by investigating the anti-sickling effect of βT87Q-globin using this sickle cell line.

The anti-sickling βT87Q-globin was developed by Pawliuk et al. by introducing the anti-sickling amino acid found at position 87 of g-globin into β-globin. Naturally occurring mutations (hereditary persistence of fetal Hb [HPFH] mutations) result in elevated (20%–30%) pancellular persistent expression of γ-globin that alleviates SCD complications, indicating ~20% βT87Q-globin (pancellular expression) would reverse the disease outcome. However, the challenge of transducing HSCs23 and the requirement for larger vectors containing erythroid-specific regulatory elements means that transduction and protein expression are variable in the bulk transplantation product, making the resulting in vivo transgene levels and clinical outcome for each patient unpredictable. Therefore, higher VCNs are targeted in the current clinical trials to enhance the success rate. Although the same vector type and process are being used for each patient, the results and outcomes are variable. Determining the therapeutic threshold for βT87Q-globin, along with HSC transduction optimization, would advance the approach and improve prospects for all patients.

We have generated an SCD cell line (sHUDEP-2) and transduced it with βT87Q-globin-encoding lentivirus to evaluate its anti-sickling activity. The results showed that the anti-sickling activity of βT87Q-globin increased in a dose-dependent manner. A VCN of 3.48 or more provided a significant amount of βT87Q-globin protein levels (81%–89%) and reduced sickling activity under low oxygen conditions. Most interestingly, βT87Q-globin expression reduced endogenous β- or β-globin expression at both RNA and protein levels in sHUDEP-2 or CD34+ derived cells, respectively. In β-thalassemia patients, unbound α-chains precipitate in RBC precursors leading to apoptosis and insufficient erythropoiesis. On the other hand, excessive β/β'-globin protein is degraded in β-thalassemia/SCD patients. Consistently, our results suggest predominantly degradation of excessive β/β'-globin, but also transcriptional regulatory mechanisms might be involved to provide globin chain balance. In the clinical setting, pre-processed Hb level measurements show a mixture of HbS and HbA (resulting from transfusion). This ex vivo model clearly demonstrates that increased βT87Q-globin levels reduce β'-globin protein levels, possibly as a result of endogenous controls on β-like globin levels. This globin expression regulation mechanism could be either through mRNA degradation or transcription control by a feedback mechanism. This observation needs to be evaluated further in other globin addition (i.e., γ-globin) approaches
for confirmation. Interestingly, early data on our clinical trial testing βT87Q lentiviral gene transfer to HSCs of SCD patients yielded similar observations of a reciprocal decrease in HbS.28

To investigate the possible transcriptomic mechanism responsible for globin regulation, we conducted RNA-seq on sHUDEP-2 and CD34+-derived cells. Although we did not find significant changes in any major β-globin expression regulating transcription factors (TFs), including GATA1, NF-E2, KLF1, and SCL,29,30 many long non-coding RNAs (lncRNA), whose roles in erythropoiesis are unknown, were differentially expressed in βT87Q-globin-transduced cells. lncRNAs are known for their role in regulating protein-coding gene activity in cis and in trans.31 In addition to TFs and regulatory RNAs, epigenetic regulators also control globin expression. β-Globin expression is controlled at the DNA level by methylation and formation of the heterochromatin structure.32 Further detailed studies are warranted to fully elucidate the exact molecular mechanism for the maintenance and control of globin chain mRNA and protein levels.

Overall, we developed an SCD cell line that can be used in ex vivo optimization studies before translation of promising approaches to animal models and clinical studies. Evaluation of βT87Q-globin over-expressing sHUDEP-2 and CD34+-derived cells demonstrated marked reduction in β/β+-globin transcripts and protein levels in sHUDEP-2 cells in a dose-dependent manner. These results provide insight into globin chain expression regulation and can be used for optimization in clinical trials involving β-globin gene transfer. Finally, these results are important in evaluating the promise of this approach for the hemoglobinopathies.

MATERIALS AND METHODS

Generation and Differentiation of the SCD Erythroid Cell Line
HUDEP-2 cells, an immortalized erythroid progenitor cell line (a gift from Drs. Yukio Nakamura and Ryo Kurita; RIKEN Tsukuba Branch, Ibaraki, Japan), were cultured and differentiated as previously described.16 In brief, cells were expanded in StemSpan II supplemented with 3 U/mL erythropoietin (EPO; AMGEN, Thousand Oaks, CA, USA), 50 ng/mL SCF (R&D Systems; Minneapolis, MN, USA), 10⁻⁶ M dexamethasone (DEX; VETone, Boise, ID, USA), and 1 μg/mL doxycycline (DOX; Sigma-Aldrich, Saint Louis, MO, USA). Cells were differentiated in Iscove’s modified Dulbecco’s media (IMDM; Sigma-Aldrich) supplemented with 50 ng/mL SCF, 10 μg/mL insulin (INS; Lilly, Indianapolis, IN, USA), 3 U/mL EPO, 400 μg/mL holo-transferrin (Sigma-Aldrich), 2 U/mL heparin (Sigma-Aldrich), 1 μg/mL DOX, and 5% human AB plasma (Rhode Island Blood Center, Providence, RI, USA) for 7 days.

For the SCD mutation introduction, the ribonucleoprotein complex (RNP) was first formed by incubation of guide RNA (200 pmol; Synthego, CA, USA) (5’-GUA ACG GGA GAC UUC UCC UC-3’) and Cas9 protein (200 pmol; UC Berkeley, Berkeley, CA, USA) at room temperature for 15 min. HUDEP-2 cells (5 × 10⁵ cells) were washed with phosphate-buffered saline (PBS) solution (Thermo Fisher Scientific, Waltham, MA, USA) twice and suspended in 20 μL SF buffer solution (#V4XC-2032, Lonza, Basel, Switzerland). The cells were mixed with RNP and single-strand donor DNA (300 pmol, 100 bases of β±-globin gene around the SCD mutation; IDT, Coralville, IA, USA). Cells were electroporated with Amaxa 4D Nucleofector X Unit (Lonza) using the manufacturer-optimized FF-120 program. The electroporated cells were incubated at room temperature for 10 min and added to pre-warmed 80 μL expansion medium before transferring into 12-well cell culture plates (Corning, NY, USA). Granulocyte colony-stimulating factor-mobilized hematopoietic cells were collected from healthy donors and perixafo-mobilized SCD patients enrolled in institutional review board-approved studies. Apheresis product underwent density gradient centrifugation, and CD34 cells were positively selected using a Miltenyi magnetic bead system. Flow cytometry was used to confirm and quantify CD34+ cell purity (>90% CD34+ and >90% viability).

Hb Electrophoresis
Hb types in differentiated cells (1–3 × 10⁶ cells) (on day 7 of differentiation for sHUDEP-2 and on day 14 for CD34+ cells) were determined using cellulose acetate membranes and alkaline buffer solution according to the manufacturer’s instructions (Helena Laboratories, Beaumont, TX, USA).

qPCR and Lentiviral Transduction of sHUDEP-2 Cells
After determining the presence of HbS in the bulk population, the culture was single-cell cloned. The clones were subjected to qPCR analysis using PCR-based genotyping (Sickle FWD4 primer: 5’-GG CAG AGC CAT CTA TTG CTT AC-3’, Sickle REV2 primer: 5’-CCA ACT TCA TCC ACG TTC ACC-3’, and Sickle probe: 5’-FAM-CTG ACT CCT GTG GAG AA-3’). A selected sHUDEP-2 clone (2 × 10⁵ cells) was pre-stimulated with fresh 1 mL expansion media overnight in a 12-well plate. The culture medium was renewed with a fresh 1 mL expansion medium containing a βT87Q-globin expressing lentiviral vector at different MOIs (0.5, 1, 2, 5, 10, and 25) and proamine (2 mg/mL; Sigma-Aldrich). The next day, transduced cells were split into two wells in 12-well plates in 1 mL fresh expansion media. Transduced cells at different MOIs were single-cell cloned, and genomic DNA from each clone was extracted using an automated DNA system following the manufacturer’s recommendation (Biomek 400; Beckman Coulter, Brea, CA, USA). VCN was determined for each clone by qPCR using the QuantStudio 6 Flex real-time PCR system (Thermo Fisher Scientific) as previously described.15 In addition, globin transcripts per α-globin was calculated as described previously.15

Flow Cytometry Analysis
Differentiated cells were characterized in terms of erythroid-specific cell surface markers using flow cytometry analysis. In brief, 1 × 10⁵ sHUDEP-2 cells were washed with PBS three times and incubated with CD36 (#555454; Becton Dickinson Biosciences, Franklin Lakes, NJ, USA), CD71 (#551374; Becton Dickinson Biosciences), and GPA (#55570; Becton Dickinson Biosciences) antibodies for 30 min at 4°C. Cells were washed with PBS and analyzed using flow cytometry.
Morphological Evaluation/Enucleation Analysis of Differentiated Cells

For morphological analysis, differentiated shUHDEP-2 cells (2 × 10⁵ cells) were centrifuged onto Shandon EZSingle Cytotunnel (Thermo Fisher Scientific) at 2,000 rpm for 2 min. Air-dried slides were stained with Wright-Giemsa and examined with light microscopy (Olympus, Tokyo, Japan) equipped with Dual-CCD camera. To determine enucleation efficiency, we stained differentiated cells with Hoechst 33258 (5 μg/mL; BD Biosciences) for 30 min at 37°C and analyzed them using LSRFortessa (Becton Dickinson).

RNA-Seq and Data Analysis

The sequencing libraries were constructed from 100 to 1,000 ng total RNA from experimental groups (n = 1) using Illumina’s TruSeq Stranded Total RNA kit with Ribo-Zero Globin (Illumina, San Diego, CA, USA) following the manufacturer’s instruction. The fragment size of RNA-seq libraries was verified using the Agilent 2100 Bioanalyzer (Agilent, Palo Alto, CA, USA), and the concentrations were determined using Qubit instrument (LifeTech, Waltham, MA, USA). Libraries were loaded onto the Illumina HiSeq 3000 for 2 × 75 bp paired-end read sequencing. Fastq files were generated using bcl2fastq software for further analysis. Quality control of paired-end reads was assessed using FastQC tools. Adaptor sequences were trimmed using Trimmomatic. Reads were aligned to the human reference genome (GRCh38) using HISAT2. FeatureCounts was used for gene-level abundance estimation using the GENCODE v.29 comprehensive gene annotations. Genes were kept in the analysis if they had raw read counts >5 in at least one sample. LImma was used to process the read counts into log₂ counts per million (log₂cpm). The log₂cpm values were normalized between samples using the trimmed mean of M values (TMM). The shUHDEP-2 and CD34⁺ samples were processed and normalized separately. Fold change ≥ 1.4 was used as a cutoff in the analysis. The datasets have been deposited in the Gene Expression Omnibus (GEO) database at the National Center for Biotechnology Information (NCBI) under the accession number GEO: GSE148529.

Statistical Analysis

The data were statistically analyzed with GraphPad Prism Software (v.6.05; GraphPad Software, USA) using one-way analysis of variance and Tukey’s post hoc test. Experiments were performed in triplicate (experimental replicates), and standard errors were represented as error bars. The p values <0.05 were considered to be statistically significant.

SUPPLEMENTAL INFORMATION

Supplemental Information can be found online at https://doi.org/10.1016/j.omtm.2020.04.013.

AUTHOR CONTRIBUTIONS

S.D., J.F.T., and N.U. designed the experiments, analyzed and interpreted data, and wrote the paper. S.D., J.J.H.-M., T.N., C.D., M.Y., and J.G. conducted cell culture and molecular experiments. B.G. and F.S. performed RNA-seq data analysis. Q.L. performed ex vivo sickling experiments.

CONFLICTS OF INTEREST

The authors declare no competing interests.
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