CRISPR/Cas9 Genome Engineering in Engraftable Human Brain-Derived Neural Stem Cells

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HIGHLIGHTS
Human neural stem cells are amenable to CRISPR-Cas9-mediated gene targeting

Truncated CD19 surface marker allows for enrichment of gene-targeted NSCs to >90%

Gene-targeted NSCs engraft, migrate, and differentiate in immunodeficient mice

GALC-engineered overexpressing NSCs cross-correct Krabbe disease fibroblasts in vitro
CRISPR/Cas9 Genome Engineering in Engraftable Human Brain-Derived Neural Stem Cells

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SUMMARY

Human neural stem cells (NSCs) offer therapeutic potential for neurodegenerative diseases, such as inherited monogenic nervous system disorders, and neural injuries. Gene editing in NSCs (GE-NSCs) could enhance their therapeutic potential. We show that NSCs are amenable to gene targeting at multiple loci using Cas9 mRNA with synthetic chemically modified guide RNAs along with DNA donor templates. Transplantation of GE-NSC into oligodendrocyte mutant shiverer-immunodeficient mice showed that GE-NSCs migrate and differentiate into astrocytes, neurons, and myelin-producing oligodendrocytes, highlighting the fact that GE-NSCs retain their NSC characteristics of self-renewal and site-specific global migration and differentiation. To show the therapeutic potential of GE-NSCs, we generated GALC lysosomal enzyme overexpressing GE-NSCs that are able to cross-correct GALC enzyme activity through the mannose-6-phosphate receptor pathway. These GE-NSCs have the potential to be an investigational cell and gene therapy for a range of neurodegenerative disorders and injuries of the central nervous system, including lysosomal storage disorders.

INTRODUCTION

Human neural stem cells (NSCs) are self-renewing, multilineage-producing cells that hold great promise for treating a large number of central nervous system (CNS) disorders. One specific NSC identified has demonstrated self-renewal at the single-cell level (Uchida et al., 2000). Based on these initial characterizations, individual NSC banks (to distinguish from freshly isolated NSCs) have been successfully established from single donated human fetal brain tissue (16–20 weeks’ gestation) by directly isolating CD133+/CD24-/lo NSCs with selective propagation in a defined serum-free culture media grown as neurospheres and cryopreserved (and therefore banked). Previous studies have shown that these NSCs are capable of proliferation, migration, and multilineage differentiation in a site-appropriate manner when they were transplanted into brains or spinal cord of immunodeficient mice, thereby recapitulating temporal development stage of NSCs from the human fetal brain (Uchida et al., 2012). No genetic modification or immortalization was required, and these banked cells continue to express NSC markers such as CD133 and Sox2 (Carpenter et al., 1999; Cummings et al., 2005; Tamaki et al., 2002, 2009; Tsukamoto et al., 2013; Uchida et al., 2000, 2012). They have biological NSC activities with multiple mechanisms of actions, providing neuroprotection, myelination, and retinal preservation via site-appropriate global migration.

Banked NSCs have been transplanted into a mouse model of lysosomal storage disorder (LSD) with PPT1 deficiency (also known as infantile neuronal ceroid lipofuscinosis [NCL] or Batten disease), and it has been demonstrated that these cells engrafted and migrated extensively, which led to reduced accumulated toxic storage material and ultimately broad neuroprotection of host cells in the hippocampus and cortex (Tamaki et al., 2009). These preclinical studies led to the first-in-human NSC transplantation for infantile and late-infantile NCLs authorized by the US Food and Drug Administration (Selden et al., 2013). Banked NSCs can also induce functional myelin in a severe demyelination model (Uchida et al., 2012), the shiverer mouse, and restore motor function in a mouse model of spinal cord injury (Cummings et al., 2005; Salazar et al., 2010). The proof-of-concept data in pre-clinical studies led to other phase I or II studies with NSCs in the hypomyelinating disease, Pelizaeus-Merzbacher disease (PMD) (Gupta et al., 2012), traumatic thoracic and cervical spinal cord injury (Levi et al., 2017), and age-related macular degeneration (clinical trial number...
These clinical studies demonstrated the safety, feasibility, long-term survival, and early evidence of efficacy of using banked NSCs in treating neurologic diseases.

One approach to increasing the therapeutic efficacy of banked NSCs would be to enhance their potential through genome engineering technologies (Porteus, 2016). Recently, transcription activator-like effector nucleases (TALENs) (Boch et al., 2009) and RNA-guided endonucleases of the CRISPR/Cas9 system (CRISPRs) have both been engineered to create site-specific DSBs in human cells. In particular, the CRISPR/Cas9 platform allows for ease, versatility, and efficacy of creating locus-specific DSBs in human cells (Jinek et al., 2012; Cong et al., 2013). The CRISPR system (derived from Streptococcus pyogenes) uses RNA-DNA Watson-Crick hybridization to identify the DNA target site to be cleaved. CRISPR deploys two components to create DSBs: (1) a 100-nucleotide (nt) RNA guide (single guide RNA [sgRNA]) of which 20 nt recognize the target site and (2) the Cas9 protein molecule that cleaves both DNA strands creating a DNA double-strand break (DSB). The resulting DSB can be resolved by one of the two highly conserved repair mechanisms, canonical non-homologous end joining (NHEJ) or homologous recombination (HR) (Kass and Jasin, 2010; Porteus, 2016). Canonical NHEJ repair functions throughout the cell cycle to correct erroneous breaks through ligation of DNA without end processing, usually resulting in small insertions or deletions (INDELs). On the contrary, HR occurs specifically during S or G2 phase of the cell cycle when an undamaged sister chromatid is available. If the cell uses HR to repair a DSB with an exogenous donor template, then precise nucleotide changes to the genome can be introduced (genome editing).

Genome editing of human neuronal-derived lineages has been mostly accomplished through embryonic stem cell (ESCs) or induced pluripotent stem cell (iPSCs) methodologies (Heidenreich and Zhang, 2016; Gaj et al., 2017). Although iPSCs allow for neuronal disease modeling, like Parkinson and Alzheimer diseases, and also functional genomic analyses during formation of complex neuronal circuits, their clinical use for CNS disorders has been limited thus far because of safety concerns (Ben-David and Benvenisty, 2011). Recently, the first CRISPR/Cas9 gene targeting was shown in NSC mammalian cell lines. This article highlighted that CRISPR/Cas9 can be used to edit safe harbor loci AAVS1 and Rosa26, biallelically knock out as well as epitope tag essential neuronal transcription factors, and engineer in glioma-initiating mutations. However, assessment of NSC function and multilineage potential in vivo as well as engineering NSCs for therapeutic value was missing (Bressan et al., 2017). The banked NSCs on the other hand have been extensively characterized in vivo and also have been used in multiple clinical trials, making genome editing of somatically derived NSCs a more feasible clinical approach for CNS disorders in the near future.

Here, we describe proof-of-concept genome editing studies in human-brain-derived propagated and banked NSCs that demonstrate the following: (1) genome editing in NSCs (GE-NSC) at three (IL2RG, HBB, CCR5) loci; (2) development of a simplified and robust cell surface marker enrichment scheme to generate a population of GE-NSCs using magnetic microbead technologies with greater than 90% purity; (3) GE-NSCs display robust engraftment and maintain biological NSC properties showing migration, differentiation, and long-term self-renewal and survival in vivo; and (4) generation of GALC lysosomal enzyme producing GE-NSCs that can cross-correct GALC enzyme activity in fibroblasts from patients affected with Krabbe disease. Collectively, these studies set the foundation for engineering and manufacturing human NSCs using genome editing to increase therapeutic potency of these cells to treat CNS disorders.

**RESULTS**

**Engineered Nuclease (CRISPR/Cas9 and TALEN)-Mediated Homologous Recombination in NSCs**

We first compared genome-editing efficiencies at the NSC “safe harbor” locus, interleukin-2 receptor gamma chain (IL2RG), in NSCs using IL2RG-specific TALEN pairs and the CRISPR/Cas9 system. The IL2RG locus is a safe harbor for transgene expression in NSCs because it is not expressed in NSCs or any NCS progeny and people with mutations in the IL2RG, although have a devastating severe combined immunodeficiency (SCID-X1) have no neurologic problems. To characterize the frequency of DSBs induced, we electroporated plasmids encoding either optimized IL2RG-specific TALEN pairs or CRISPR components (Cas9 and sgRNA), cultured cells for 7 days, isolated genomic DNA (gDNA), and then analyzed INDEL frequencies. The CRISPR system generated an average of 15% INDELs compared with 5% using the TALEN (Figure S1A, Related to Figure 1). We next compared mRNA delivery of IL2RG TALEN pairs to mRNA delivery of Cas9 along with chemically modified sgRNAs (“All RNA”). Electroporating NSCs...
with the All RNA CRISPR system induced 5-fold more INDELs than mRNA delivery of TALENs (Figure S1B, related to Figure 1). As both engineered nuclease systems were inducing DSBs (albeit at different frequencies), we targeted NSCs for HR with a plasmid encoding a homologous 

IL2RG cDNA-UbC-GFP donor cassette (Figure 1A), or a plasmid that introduced single nucleotide changes. Targeted NSCs were cultured for at least 30 days to allow episomal donor expression to dilute out such that fluorescence-activated cell sorting analysis would measure the frequency of cells with cassettes stably integrated by HR. Plasmid-based delivery of both engineered nuclease platforms mediated stable GFP expression in an average of 3.5% of NSCs (Figure S1C, related to Figure 1), whereas by using restriction fragment-length polymorphism to analyze the frequency of targeted alleles, the CRISPR system consistently modified more IL2RG alleles (Figure S1D, related to Figure 1). These data suggested that the CRISPR system was superior to TALENs, and therefore, was the nuclease used for the remainder of the studies described. These data also show that

**Figure 1. CRISPR/Cas9-Mediated Homologous Recombination (HR) in NSCs**

(A) The IL2RG locus was targeted for homologous recombination (HR) by creating double-strand breaks (DSBs) using Cas9 (scissors) and supplying a homologous donor template (with flanking 800-bp arms around the transgene to be inserted). Following HR, IL2RG cDNA is knocked in to the endogenous start codon with a UbC promoter driving GFP downstream to assess efficiencies of HR in human NSCs. IL2RG exons are shown in black boxes.

(B) NSCs were electroporated with 1-μg plasmid (px330) encoding Cas9 and sgRNA specific for IL2RG, HBB, or CCR5. For “All RNA” experiments, Cas9 was delivered as mRNA (5 μg), and the locus-specific sgRNAs (10 μg) were delivered with terminal chemical modifications comprising 2’-O-methyl 3’phosphorothioate (MS) or 2’-O-methyl 3’tioPACE (MSP). Following 7 days in culture, gDNA was harvested, PCR was carried out to amplify the region of interest, and INDELs were calculated using the Tracking of Indels by Decomposition (TIDE) software. (N = 2–14), **p < 0.01, Student’s t test.

(C) NSCs were nucleofected as described above with either 2 μg HR donor or 2 μg HR and locus-specific CRISPR components. Locus-specific HR donor plasmids have at least 400 bp of homology flanking UbC-GFP to assess HR efficiencies. Cells were grown for at least 30 days until episomal HR donor was lost by dilution during proliferation to confirm stable expression of integrated cassettes into the IL2RG, HBB, and CCR5 loci. Representative fluorescence-associated cell sorting plots are shown, which highlight stable integration of HR donors with site-specific nucleases.

(D) HR frequencies obtained from all experiments targeting IL2RG, HBB, and CCR5 carried out as above. (N = 3–7), *p < 0.05, Student’s t test.

(E) gDNA was harvested from experimental groups at day 30 post-culture to evaluate site-specific integration of HR donors on the DNA level. Digital droplet PCR (ddPCR) was used to quantify allelic integration of HR donors, (N = 3–4). Agarose gel electrophoresis of in-out PCR products confirms on-target integration of HR donors in the presence of a site-specific nuclease. Data are represented as mean ± SEM.
the IL2RG locus is an effective safe harbor in NSCs not only because the gene plays no functional role in these cells but also because it is permissive for high-level transgene expression when a donor cassette using a ubiquitous promoter is inserted into the locus, while presumably not affecting the expression of neighboring NSC genes.

We next tested whether the CRISPR system could edit two other NSC “safe harbor” loci, the chemokine (C-C motif) receptor 5 (CCR5) and β-globin (HBB) genes. Both genes are not expressed in NSCs or their progeny, and people with mutations in these genes have no neurologic consequences from those mutations and thus qualify as safe harbors in NSCs. Plasmids encoding Cas9 and sgRNAs specific to IL2RG, HBB, and CCR5 were electroporated into NSCs, and 7 days later, gDNA was extracted and alleles were analyzed for INDEL frequencies. The CRISPR-Cas9 system induced an average of 12%, 6%, and 36% INDELs at the IL2RG, HBB, and CCR5 loci, respectively (Figure 1B). Consistent with our previous reports, delivery of Cas9 as mRNA and chemically modified sgRNAs induced more INDELs than plasmid delivery at all three loci tested (Figure 1B). To see if DSB formation promoted HR at all three loci, NSCs were co-electroporated with homologous DNA donor templates encoding GFP, and corresponding CRISPR components delivered as plasmid or All RNA. Our results showed that all three loci were amenable to HR in NSCs using plasmid or All RNA delivery of CRISPR-Cas9 components (Figures 1C and S2, related to Figure 1). Targeting NSCs at IL2RG, HBB, and CCR5 loci with plasmid delivery of CRISPR resulted in an average of 2.8%, 4.1%, and 2.4% GFP+ NSCs after at least 30 days in culture, respectively (Figure 1D). Interestingly, All RNA delivery of IL2RG CRISPR resulted in 2-fold more GFP+ NSCs than plasmid delivery at all three loci tested (Figure 1B). To see if DSB formation promoted HR at all three loci, NSCs were co-electroporated with homologous DNA donor templates encoding GFP, and corresponding CRISPR components delivered as plasmid or All RNA. Our results showed that all three loci were amenable to HR in NSCs using plasmid or All RNA delivery of CRISPR-Cas9 components (Figures 1C and S2, related to Figure 1). Targeting NSCs at IL2RG, HBB, and CCR5 loci with plasmid delivery of CRISPR resulted in an average of 2.8%, 4.1%, and 2.4% GFP+ NSCs after at least 30 days in culture, respectively (Figure 1D). Interestingly, All RNA delivery of IL2RG CRISPR resulted in 2-fold more GFP+ NSCs than plasmid delivery. To determine if HR occurred at the intended loci, gDNA was analyzed for on-target integration of the donor construct by in-out digital droplet PCR (ddPCR) (where one primer binds outside the homology arm of the donor construct and the other primer binds inside the donor cassette), thus only on-target integration events can be PCR quantified. PCR amplification was not detected with mock or donor only samples, whereas on-target integration was evident when NSCs were co-electroporated with CRISPR and a homologous donor, confirming HR at the intended locus (Figure 1E). As seen before, All RNA delivery of IL2RG CRISPR increased the frequency of targeted alleles, whereas HBB and CCR5 All RNA CRISPR delivery mediated fewer on-target integrations. Of note, we report an under-representation of on-target alleles compared with GFP+ cells for targeting CCR5 and HBB, which may be due to non-target integrations of the donor-plasmid DNA, incomplete HR that interrupted ddPCR primer or probe sequences in the bovine growth hormone polyadenylation (BGH) poly(A) region, or the fact that there is still some episomal DNA expressing GFP after 30 days in culture in CRISPR-treated human NSCs. Because Cas9 can create DSBs at unintended genomic locations (Wu et al., 2014), we investigated off-target DSB activity at three predicted IL2RG off-target sites that we have previously assayed by next-generation sequencing in K562 cells (Hendel et al., 2015). We detected 28% DSB activity at IL2RG, whereas less than 0.7% cleavage was seen at any of the off-target sites investigated (Table 1). Collectively, these studies demonstrated that CRISPR-Cas9 can mediate on-target HR at multiple “safe harbor” loci in NSCs with minimal off-target cleavage. Thus these data establish that all three loci can be effective safe harbors in human NSCs as they support transgene expression from a cassette using an endogenous ubiquitous promoter. These data also provide further evidence that HR-mediated genome editing can be active at loci that are not expressed (Cameron et al., 2017) and that even though these loci are not naturally transcriptionally active in NSCs, they can still support transgene expression when an expression cassette is inserted into them. The level of transgene expression at the HBB locus is lower than at the other two loci, which might be used to tailor expression levels in certain applications.

| Locus       | Unselected | CD8 Selected | CD8 Selected | tCD19 Selected |
|-------------|------------|--------------|--------------|----------------|
| IL2RG       | 28.35      | 75.85        | 59.11        | 71.27          |
| Off-target 1 | 0          | 0            | 0.01         | 0              |
| Off-target 2 | 0          | 0.28         | 0.07         | 0              |
| Off-target 3 | 0          | 0            | 0            | 0.62           |

Table 1. Low Off-Target Activity of IL2RG CRISPR/Cas9 System in Human NSCs
Targeted deep sequencing of IL2RG alleles and three previously predicted off-target sites was performed on unedited NSCs and GE-NSCs. Each column represents an independent targeting experiment from GE-NSCs that were unselected, CD8 selected, or tCD19 selected. Off-targets were previously predicted using bioinformatic tools (Hendel et al., 2015). All numbers are subtracted from background INDEL frequencies in an unedited NSC sample.
Enrichment of Genome-Edited NSCs (GE-NSCs) by Magnetic-Activated Cell Sorting (MACS) Selection

Enrichment and expansion of genome-edited NSCs (GE-NSCs) to clinically relevant transplantable cell numbers would greatly increase their therapeutic potential. We devised an enrichment paradigm by establishing expression of the alpha chain of the CD8 protein dimer (CD8α) following successful HR at IL2RG coupled with a one-step magnetic-activated cell sorting (MACS) enrichment. CD8 complex requires dimerization of alpha and beta chains to transduce intercellular signals in cytotoxic T cells. Of note, expression of CD8α chain only has been reported not to be functional and should not alter NSC biologic function. In contrast, the truncated nerve growth factor (NGF) receptor might be predicted to perturb the natural signaling pathways of the brain by serving to “steal” NGF from cells that need NGF for their function. Thus the CD8 cell surface molecule on these cells can be used as a biologically inert cell surface marker to enrich for genetically modified NSCs. We therefore created an IL2RG HR donor construct (IL2RG cDNA-UbC-GFP-T2A-CD8α) that upon successful targeting would express a multigene mRNA, but two distinct GFP and CD8α proteins. NSCs were electroporated with a plasmid encoding Cas9 and sgRNA specific to IL2RG and also the CD8α HR donor cassette described above. Six cell passages post-electroporation, 4.7% of NSCs were stably expressing GFP (Figure S3A, left, related to Figure 2), similar to what we found with the GFP-only construct (Figure 1C). Then NSCs were subjected to a one-step CD8α magnetic bead enrichment protocol that resulted in >90% of cells expressing GFP (Figure S3A, right, related to Figure 2), and these cells were shown to expand and continually express GFP for multiple passages post-selection, confirming they were a population of GE-NSCs (Figure 2A).

Figure 2. Enrichment of Genome-Edited NSCs (GE-NSCs) by Magnetic-Activated Cell Sorting (MACS) Selection

A bicistronic IL2RG HR cassette was created that separates GFP from CD8 or truncated CD19 (tCD19) via a T2A peptide motif to allow robust magnetic bead enrichment of IL2RG-targeted NSCs. NSCs were nucleofected with 2 μg HR donor and 1 μg plasmid encoding Cas9 and sgRNA. Cells were grown for 30 days to allow episomal HR donor to dilute out during proliferation of NSCs.

(A) GE-NSCs were cultured for nine passages post-electroporation while being analyzed for GFP expression at every cell passage. Cells were selected using CD8 microbead technologies at passage 6 post-targeting.

(B) GE-NSCs were cultured for nine passages while being analyzed for GFP expression.

(C) Frequencies of GE-NSCs using the bicistronic donor cassettes before (targeted) and after MACS selection (enriched), N = 3, Data are represented as mean ± SEM.

(D–F) Expanded/MACS-enriched GE-NSCs with bicistronic GFP-T2a-tCD19 cassette were enzymatically dissociated into a single cell suspension and then stained for the cell surface markers, CD133 (Phycoerythrin [PE]) and CD19 (Allophycocyanin [APC]), and analyzed by FACS; (D) CD19 versus GFP and (E) CD19 versus CD133. (F) Cells were stained for CD19 (APC), permeabilized, fixed, and then stained again for SOX2 (PE).

Enrichment of Genome-Edited NSCs (GE-NSCs) by Magnetic-Activated Cell Sorting (MACS) Selection

In addition to CD8α, we also evaluated CD19 cell surface protein for our MACS-based enrichment protocol. Unlike the CD8 complex, CD19 is a single-chain molecule, and functional CD19 has cytoplasmic phosphor-ylated tyrosine domain, which transduces a signal (Depoil et al., 2008). Therefore we created a truncated form of the CD19 cell surface protein, where the intracellular signaling domain is removed, thus making CD19 signaling inert and solely serving as a cell surface marker for enrichment. In addition, even full-length
CD19 has no known role in neurogenesis or gliogenesis and is not known to be expressed in any neural lineage. We therefore created an IL2RG HR donor construct (IL2RG cDNA-UbC-GFP-T2A-tCD19) and targeted NSCs for HR at the IL2RG locus. At passage four post-electroporation, 3.8% of NSCs were stably expressing GFP and were then subjected to MACS-based enrichment of tCD19 (Figure S3B, left, related to Figure 2). tCD19 selection resulted in enrichment and expansion of >90% of GE-NSCs, similar to CD8 selection of GE-NSCs (Figures 2A–2C). Using this enrichment methodology, it is also possible to select for cells with biallelic integration using two DNA donor molecules, each with a different fluorochrome (Bak et al., 2017). To test this in NSCs, we targeted the CCR5 locus with both UbC-GFP and UbC-mCherry-T2A-tCD19 constructs. Targeting with both constructs resulted in 0.19% cells with targeted integration at both alleles, and CD19 MACS enrichment resulted in an ~100-fold increase (12%) of biallelically targeted NSCs (Figure S4, related to Figure 2). Because we are enriching cells with HR events, we next wanted to investigate whether we were also selecting GE-NSCs with a higher frequency of off-target INDELs. Although we observed a 3-fold enrichment of IL2RG INDELs (because the banked NSCs are female with two X chromosomes), off-target INDELs were all less than 1% in CD8- and tCD19-selected GE-NSCs (Table 1).

To determine if GE-NSCs maintained their NSC characteristics, we enriched IL2RG-targeted NSCs to over 95% (Figure 2D) via the tCD19 selection scheme and then analyzed the expression of the two quintessential NSC markers: the cell surface gene CD133 (Figure 2E) and the intracellular NSC-maintaining transcription factor SOX2 (Figure 2F) (Hook et al., 2011). Following four weeks of expansion post-enrichment, greater than 95% of NSCs were CD133/CD19/GFP/SOX2 positive, highlighting that targeted neurospheres may fully possess NSC potential. These data provide two separate examples of the feasibility of a simplified one-step MACS-based protocol for enriching and then expanding >90% of GE-NSCs for transplantation.

Although our MACS-based enrichment strategy consistently resulted in purification to >90% of GE-NSCs, we also tested the recombinant adeno-associated virus serotype six (rAAV6) homologous donor template platform that has been shown to work well in primary cells (Dever et al., 2016; Sather et al., 2015; Wang et al., 2015). We electroporated Cas9 mRNA and HBB MSP sgRNAs into NSCs and then immediately transduced them with rAAV6 (Bak et al., 2018) carrying a UbC-GFP cassette with arms of homology for HBB. Using vector genomes per cell of 10,000, 100,000, and 500,000, we were able to achieve mean allelic targeting frequencies of 1.26%, 9.93%, and 12.57%, respectively (Figure S5, related to Figure 2). These results show that AAV6 is a highly active homologous donor in human NSCs that warrants future investigation. Given the enrichment to >90% purity using the one-step magnetic bead purification from using the plasmid donor, we pursued further in vivo analysis using these GE-NSCs.

**GE-NSCs Migrate and Differentiate into Neurons, Astrocytes, and Oligodendrocytes In Vivo**

While we have shown that we can target NSCs for HR using the CRISPR/Cas9 system and after engineering they retained marker expression consistent with NSC function, we investigated if GE-NSCs still retain their NSC biological activity as previously shown for non-edited NSCs (Tsukamoto et al., 2013). To test this hypothesis, we targeted the IL2RG locus in NSCs with a GFP-T2A-CD8α cassette (Figure 1A), expanded GE-NSCs, and then transplanted them into immunodeficient mice. CD8α-purified GE-NSCs were transplanted bilaterally into the subventricular zone (SVZ) of neonatal shiverer-immunodeficient (shi/shi-id) mice, and then after 12 weeks, serial forebrain sections from the same animal were analyzed for engraftment of human modified cells by immunohistochemistry (Figure 3A). Immunoperoxidase staining with the human-specific mAb SC121 (Figure 3A top) and GFP (Figure 3A bottom) detected engraftment of GE-NSCs in the cortex and corpus callosum and importantly, showed migration of cells from the SVZ to the olfactory bulb along the rostral migratory stream (RMS) (Figure 3A). GFP expression was very similar to human-specific SC121 expression, indicating that GE-NSCs can engraft and migrate in the SVZ/olfactory system, comparable to non-edited NSCs reported previously (Tsukamoto et al., 2013).

The hallmark of human NSCs is their ability to self-renew while differentiating into neuronal, astrocytic, and oligodendrocytic lineages in long-term xenograft transplantation studies. GFP+ cells were detected in the hippocampus in shi/+ -id mice 24 weeks after transplant demonstrating long-term survival and migration of the GE-NSCs. In addition, SOX2-expressing human cells were detected in the subgranular layer of the dentate gyrus of the hippocampus, suggesting the maintenance of the human NSC in the neurogenic niche (Figure 3B arrows). Confocal microscopy on brain sections were performed to confirm the ability of the GE-NSC to differentiate into astrocytic and neuronal cells using the markers, glial fibrillary acidic protein (GFAP), and NeuN.
We observed GFP-positive cells costained with GFAP in the corpus callosum or white matter bundle of the striatum (Figure 3C). Furthermore, GE-NSCs migrated into the RMS and differentiated into DCX+ neuronal lineage in the olfactory bulb (Figure 3D), revealing that migration and differentiation of GE-NSC into astrocytic and neuronal lineages occurs as expected.

The differentiation potential of GE-NSCs into oligodendrocytes was evaluated using the oligodendrocyte mutant shi/shi-id as described previously (Uchida et al., 2012). The shi/shi-id mouse has a major deletion in myelin basic protein (MBP) gene, rendering oligodendrocytes deficient at producing myelin in the white matter tracks. Eight weeks post-transplant, GE-NSCs migrated within the white matter tract of the cerebellum (Figure 3E) and expressed the knocked in transgene GFP (Figure 3F). Immunofluorescence staining revealed that the engrafted cells in the white matter co-expressed GFP and MBP (Figure 3G). In a higher magnification of a different mouse brain, we observed that human GFP+ cells extended their processes and MBP expression was localized at the tip of their process, indicative of myelinating oligodendrocytes (GFAP) and doublecortin (DCX), respectively. We observed GFP-positive cells costained with GFAP in the corpus callosum or white matter bundle of the striatum (Figure 3C). Furthermore, GE-NSCs migrated into the RMS and differentiated into DCX+ neuronal lineage in the olfactory bulb (Figure 3D), revealing that migration and differentiation of GE-NSC into astrocytic and neuronal lineages occurs as expected.

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Thus, GE-NSCs maintained the potential to engraft, migrate, self-renew, and differentiate into the three CNS lineages in a site-appropriate manner as we have seen for unedited NSCs (Uchida et al., 2000, 2012; Tamaki et al., 2009). In addition, the unedited NSCs have not demonstrated tumor formation in mouse or humans and the edited NSCs similarly did not generate tumors following transplantation into mice.

**GALC Lysosomal Enzyme-Overexpressing NSCs Cross-Correct Deficient Enzyme Activity in Krabbe Disease Cells**

LSDs are a group of more than 50 inherited monogenic metabolic disorders with accumulated toxic materials that primary result from the deficiency of lysosomal enzyme (Parenti et al., 2015). Among them, globoid cell leukodystrophy, or Krabbe disease, is a type of LSD that mainly manifests in the central and peripheral nervous systems due to the loss of the enzyme galactosylceramidase (GALC), which causes death of myelin-producing oligodendrocytes and Schwann cells, respectively. NSCs have been transplanted into the brains of patients with NCL to deliver the missing enzyme (Selden et al., 2013) and in patients with PMD to provide myelin-producing cells (Gupta et al., 2012). As we have shown that GE-NSCs produced myelin comparable to unedited NSCs, we reasoned that GE-NSCs overexpressing GALC protein would be superior in terms of producing enough GALC to enable cross-correction of damaged myelin-producing cells. Moreover, these cells could also replace already lost oligodendrocytes with “normal”-GALC expressing oligodendrocytes. Therefore we created an HR donor to knock in GALC (IL2RG cDNA-UbC-GALC-T2A-tCD19) into the IL2RG locus and also a tCD19 selection cassette to enable robust MACS-based enrichment (Figure 4A). NSCs were electroporated with the “All RNA” IL2RG CRISPR/Cas9 platform and the GALC HR donor with a mean of 2.95% CD19-positive NSCs after episomal donor DNA was diluted out at least four passages post-targeting (Figure 4B). Consistently, tCD19 MACS beads enriched to >90% of GALC GE-NSCs (Figure 4B) that accordingly maintained neuronal stem cell phenotypes (Figure S6A, related to Figure 4) and enrichment of safe harbor IL2RG on-target integration events (Figure S6B, Related to Figure 4).

To confirm that GALC NSCs were overexpressing functional enzyme, we performed an in vitro GALC enzyme assay that was validated by comparing GALC knockout Krabbe disease fibroblasts (Figure S6C, related to Figure 4) with GALC K562 cells that were generated by knocking UbC-GALC-T2A-GFP into the IL2RG locus (Figure S6D, related to Figure 4). Accordingly, GALC GE-NSCs expressed 10-fold more GALC enzyme than unedited NSCs in whole protein lysates and 3-fold more in the cultured media (Figure 4C, left). As GALC GE-NSCs are overexpressing active GALC enzyme and releasing it into the culture media, we performed co-culture experiments with previously characterized Krabbe disease fibroblasts (Luzi et al., 1995; Rafi et al., 1995) to ascertain if they could be cross-corrected for GALC enzyme activity by the GALC NSCs. After 7 days (1 passage) and 14 days (2 passages) of trans-well-mediated co-culturing of GALC NSCs with Krabbe disease fibroblasts, GALC enzyme activity was increased 18- and 34-folds, respectively (Figure 4C, right). These results suggest that active GALC enzyme is accumulating linearly as the co-culture time increases. To confirm that GALC cross-correction was mediated through the essential lysosomal mannose-6-phosphate (M6P) receptor pathway (Tamaki et al., 2009), we cultured with M6P to antagonize the enzyme uptake in Krabbe disease fibroblasts. We were able to inhibit 73% of GALC enzyme uptake with M6P added to the culture media every 24–48 h (Figure 4D), suggesting that the cross-correction of GALC enzyme activity to Krabbe disease cells is via the M6P receptor lysosomal pathway.

We also confirmed that GALC GE-NSCs retained their NSC biological characteristics by transplanting GALC GE-NSCs into the cerebellum of juvenile Shi-id homozygous mice. Immunohistochemical analyses showed engraftment via SC121 marker expression and myelin expression via MBP expression of GALC GE-NSCs (Figure S7, related to Figure 4). These data combined with the fact that GALC-NSCs maintain neuronal stem cell markers (Figure S6A, related to Figure 4) highly suggest that GALC GE-NSCs retain their NSC biological characteristics and may be of therapeutic potential, which warrants future safety and efficacy investigation in relevant mouse models for the treatment of Krabbe disease and other demyelinating CNS disorders.

**DISCUSSION**

We have outlined a genome editing methodology in NSCs using the CRISPR/Cas9 platform. In brief, we demonstrate (1) genome editing by HR at multiple NSC safe harbor loci; (2) an enrichment strategy for selecting >90% of genome-edited NSCs (GE-NSCs) by using signaling-incompetent cell surface markers and microbead technologies; (3) that GE-NSCs retain their NSC characteristics in vivo by showing proliferation
in the hippocampal stem cell niche, migration along the rostral migratory stream, and differentiation into neurons and astrocytes as well as oligodendrocytes that produce MBP in all brain regions investigated; and (4) generation of GE-NSCs overexpressing the GALC enzyme that cross-correct GALC enzyme activity in Krabbe disease cells, and engraft and produce myelin in vivo. Overall, these data warrant more comprehensive safety and efficacy studies for the investigational GE-NSCs cell products that could be used as a therapeutic cell product for CNS disorders and also could be leveraged to study human NSC biology.

Most CRISPR/Cas9 applications in neuroscience have used NHEJ-based editing of (1) mouse brains in vivo and (2) human iPSCs and ESCs in vitro, and although these advances have helped better understand neuronal networks, there are limitations of these systems, such as nuclease delivery and restricting editing to a specific cell type (Heidenreich and Zhang, 2016). Although lentiviral-mediated gene transfer for NSCs has been shown to efficiently deliver transgenes in vitro and in vivo (Jandial et al., 2008), the concerns of insertional mutagenesis always remain high. An alternative approach is to precisely edit the locus of interest by homologous recombination directly in human NSCs that possess in vitro and in vivo multilineage potential that can be expanded and created into cell banks to provide an allogeneic product for transplantation. Because NSCs maintain trilineage potential (oligodendrocytes, neurons, astrocytes) in vitro and

Figure 4. GALC Overexpressing NSCs Cross-correct Krabbe Disease Fibroblasts through Mannose-6-phosphate Receptor Pathway
(A) The IL2RG locus was targeted for homologous recombination (HR) by creating double-strand breaks (DSBs) using Cas9 (scissors) and supplying a homologous donor template (with flanking 800-bp arms around the transgene to be inserted). Following HR, IL2RG cDNA is knocked in to the endogenous start codon followed by a UbC-driven cassette with GALC-T2A-tCD19 for overexpression of GalC enzyme and enrichment of targeted cells using CD19 MACS selection.
(B) (Left) NSCs were nucleofected as described previously with Cas9 mRNA, MSP L2RG sgRNA, and the GALC-T2A-tCD19 HR donor construct. Representative fluorescence-activated cell sorting plots highlight stably integrated GALC construct as measured by tCD19 expression at 30 days post-nucleofection before (top) and after (bottom) enrichment of tCD19+ cells (right). (Right) Frequencies of GALC-NSCs before (targeted) and after MACS selection (post enriched), N = 3–6.
(C) (Left) GALC enzyme assay was performed on culture media and cellular protein lysates from unedited NSCs and GALC GE-NSCs (two different NSC cell banks). GALC enzyme activity is presented as nmol/mg/17 h (normalized to unedited HuCNS-SCs). N = 1–4, two independent biological experiments. (Right) Krabbe fibroblasts (cell lines #6806 and #8304) were either cultured alone (no NSCs) or co-cultured with GALC GE-NSCs for 7 days (P1) or 14 days (P2). Krabbe fibroblasts were then harvested for GALC enzyme assay. GALC activity is presented as nmol/mg/17 h, N = 4–6, three independent biological experiments **p < 0.01, Student’s t test.
(D) GALC cross-correction experiments were performed as described in Figure 4C, except mannose-6-phosphate (M6P) was added to the culture media every 24–48 h (2.5 mM final concentrations) for 7 days total. Krabbe fibroblasts were harvested, protein was isolated, and GALC activity was carried out. N = 2, two independent biological experiments, *p < 0.0143, Mann-Whitney test. Data are represented as mean ± SEM.
CNS site-specific differentiation and migration in vivo, NSCs are an ideal candidate cell population to genetically engineer to explore NSC gene function programs, as banked NSCs migrate globally to facilitate neuroprotection by supplying missing lysosomal enzyme or providing neurotrophic factors. Our data show that it is possible to target NSCs for HR at multiple loci and enrich and expand genome-edited NSCs, and more importantly, our data unequivocally demonstrate that GE-NSCs maintain hallmark NSC functions: stem cell marker expression, lineage commitment, and migration in vivo. Therefore, our methodology sets the framework to allow investigators to interrogate CNS gene-cell functions at any locus of interest and also to lineage trace NSCs in vivo.

Banked NSCs have already been tested in preclinical models as well as have been used as an investigational cellular product in multiple phase I or II clinical trials for difficult-to-treat CNS disorders, such as retinal degeneration (McGill et al., 2012; Cuenca et al., 2013), spinal cord injuries (Cummings et al., 2005; Salazar et al., 2010), rare myelin disorders (Uchida et al., 2012), and LSDs (Tamaki et al., 2009). Although the outcomes of these trials were encouraging, they also highlighted the potential to facilitate improvements to the cellular product (Tsukamoto et al., 2013). Krabbe disease is an LSD that results from the loss of GALC activity in myelin-producing cells, and as NSCs already promote oligogenesis (Uchida et al., 2012), we reasoned that constitutively overexpressing GALC through HR in NSC safe harbor loci would improve their therapeutic potential. Accordingly, our data highlight that GALK-NSCs secrete enough active GALC enzyme to be taken up by Krabbe disease cells through the M6P receptor pathway and rescue deficiencies in enzyme levels. Furthermore, our current studies also show that GALK GE-NSCs expresses essential neuronal stem cell markers and are able to differentiate into oligodendrocytes (and produce myelin) in vivo. The fact that GALK-NSCs can not only secrete enough enzyme but also can differentiate into oligodendrocytes is critical for their therapeutic activity. Now, the next logical set of future studies would be to investigate whether GALC GE-NSCs that overexpress GALC enzyme in vivo have the ability to cross-correct dysfunctional host oligodendrocytes in a Krabbe mouse model and also provide donor-derived oligodendrocytes to repair hypomyelination. In terms of future experiments, our methodology also allows us to overexpress any protein of interest, and future studies will interrogate whether overexpression of neurotrophic factors, such as brain-derived neurotrophic factor, can improve neuronal survival in neurodegenerative diseases like Alzheimer, Parkinson, and Huntington models.

In conclusion, we have set the framework for CRISPR/Cas9 genome editing by HR in human-brain-derived multipotent NSCs that will advance the field of neuroscience and treatment of CNS disorders.

Limitations of the Study
We demonstrate that CRISPR-Cas9 gene-targeted human NSCs can engraft, migrate, and differentiate into CNS cells in a site-specific manner in the mouse brain. We also show that GALC-expressing NSCs can be engineered to secrete lysosomal enzymes that are able to cross-correct Krabbe fibroblasts in vitro. The limitation of this study is that we have not definitively shown in vivo correction of a Krabbe disease mouse model, such as the twitcher mouse model. Additional experiments are needed to further provide preclinical experimental proof of the feasibility and efficacy of using GE-NSCs as a treatment for LSDs.

METHODS
All methods can be found in the accompanying Transparent Methods supplemental file.

SUPPLEMENTAL INFORMATION
Supplemental Information can be found online at https://doi.org/10.1016/j.isci.2019.04.036.

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AUTHOR CONTRIBUTIONS
D.P.D and N.U performed most of the experiments. S.G.S performed the GALC enzyme assays and cross-correction experiments. J.C. performed all ddPCR analyses. E.J.K. and J.T.C. performed early nucleofection optimization experiments and gene targeting experiments. R.O.B. performed gene-editing experiments related to CCR5. R.M.M. performed the rAAV6 experiments. Y.T. and M.D. performed in vivo transplantations as well as immunohistochemistry. J.A.B., K.A.J., and G.B. analyzed off-target MiSeq libraries. N.G.-O. directed experiments performed by S.G.S. D.P.D wrote the manuscript with help from all the authors. M.H.P. and N.U equally directed the research.

DECLARATION OF INTERESTS
M.H.P. is a consultant and has equity interest in CRISPR Tx, but CRISPR Tx had no input into the design, execution, interpretation, or publication of the results herein. N.U., A.T., Y.T., and M.D. were former employees of StemCell, Inc. ReGen Med Division, BOCO Silicon Valley, acquired the intellectual properties and technologies from StemCells Inc. N.U. and A.T. are current employees of ReGen Med Division, BOCO Silicon Valley. There is a patent pending, US15/487,309, as a result of this work.

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Supplemental Information

CRISPR/Cas9 Genome Engineering in Engraftable Human Brain-Derived Neural Stem Cells

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Supplemental Figure 1. The CRISPR system is highly active in human NSCs. A) 500,000 NSCs were electroporated (plasmid delivery) with either the CRISPR/Cas9 system or TALEN pairs that were designed to recognize the human IL2RG locus. Seven days post targeting, gDNA was harvested, IL2RG alleles were amplified by PCR with primers that overlapped the cut site, and TIDE was run to analyze INDEL frequencies. (N= 4-7), * p < 0.05, Student’s T-test. B) NSCs were electroporated as described above with Cas9 mRNA and MSP sgRNA or TALEN pairs delivered as mRNA, and then IL2RG alleles were analyzed by TIDE software for INDELs. (N= 2-5), * p < 0.05, Student’s T-test. C) NSCs were electroporated with IL2RG engineered nucleases along with UbC-GFP donor templates with IL2RG homology arms. 30 days post-targeting, cells were harvested for FACS GFP analyses. (N= 4-6). D) NSCs were electroporated with IL2RG engineered nucleases along with an IL2RG homologous donor template intended to introduce a AflII restriction site following homologous recombination. 7 days post-targeting, gDNA was harvested, then IL2RG alleles were amplified to produce a 1.6kb product. Amplified alleles were digested with AflII and run a PAGE gel. The number of HR alleles were calculated as follows: ((digested alleles/ digested alleles) + undigested alleles). (N= 4-6), Student’s T-test. Data are represented as mean +/- SEM.
Supplemental Figure 2. Schematic of targeted integration into HBB and CCR5 loci. A) The HBB locus was targeted by creating a DSB in exon 1 via Cas9 (scissors) and supplying a UbC-GFP homologous donor template. Alleles with integrations were identified by PCR (881bp) using an In (black) - Out (red) primer set. B) The CCR5 locus was targeted in exon 3 as described above. PCR identified integrated (1200bp) alleles using In (black) - Out (red). Loading control for all In-Out PCRs was the wildtype CCR5 allele.
Supplemental Figure 3. Representative FACS images showing MACS enrichment of IL2RG targeted NSCs. A bicistronic $IL2RG$ HR cassette was created that separate GFP from CD8α or truncated CD19 (tCD19) via a T2A peptide motif to allow robust magnetic bead enrichment of $IL2RG$-targeted NSCs. NSCs were nucleofected with 2ug HR donor and 1ug plasmid encoding Cas9 and sgRNA. Cells were grown for 30 days to allow episomal HR donor to dilute out during proliferation of NSCs. A) Representative FACS plots show a population of NSCs with stable integration of the bicistronic GFP-T2A-CD8 cassette before enrichment at day 30 after electroporation (left), and 30 days after enrichment using Magnetic Activated Cell Sorting (MACS) CD8 Microbead technologies (right). B) Representative FACS plots showing a population of NSCs with stable integration of the bicistronic GFP-T2A-tCD19 cassette before enrichment at day 30 after electroporation (left), and a population enriched of genome-modified NSCs using MACS CD19 Microbead technologies (right).
Supplemental Figure 4. Enrichment of biallelic CCR5-targeted NSCs. A) Human NSCs were targeted at CCR5 with a UbC-GFP cassette (top middle), with a UbC-mCherry-T2A-tCD19 cassette (top right), or with both constructs (bottom middle). CD19-based MACS enrichment results in greater than a 10 fold increase in biallelically edited CCR5 NSCs as well as monoallelic CCR5 edited cells. B) Quantification of the percent of GFP targeted, tCD19 targeted or biallelically targeted NSCs from the experiment shown in A.
Supplemental Figure 5: Recombinant adeno-associated virus serotype six (AAV6) is a suitable homologous donor template in human NSCs. 500,000 human NSCs were electroporated with Cas9 mRNA and HBB-specific MSP sgRNAs and then cells were immediately transduced with rAAV6 carrying UbC-GFP with arms of homology for HBB (centered on the sgRNA cut site). A) Representative ddPCR images show the increased number of HBB targeted allele positive droplets when the number of AAV6 vector genomes per cell is increased. The aggregate data is shown in B below. The reference allele amplified is CCR2. B) In three independent experiments, NSCs were HBB targeted with a UbC-GFP AAV6 donor (at three different VGs/cell), cells were harvested 7 days later and then gDNA was harvested and ddPCR was carried out to determine the frequency of HBB integrated alleles. Using AAV6 VGs/cell of 10,000, 100,000 and 500,000, mean integration allele frequencies were 1.26%, 9.93%, and 12.57%, respectively. The lines pair VG dose responses in the same experiment.
Supplemental Figure 6: GALC-NSCs maintain neuronal stem cell markers and establishment of GALC enzyme assay.  A) Enriched GALC-NSCs were analyzed for editing via tCD19 and stem cell markers via CD133, SOX2 and SOX1. 70% of GALC-NSCs were targeted and express quintessential neuronal stem cells markers.  B) 500,000 NSCs were mock electroporated, electroporated with DNA donor only, or electroporated with GALC-T2A-tCD19 DNA donor and ALL RNA Il2RG CRISPR components. After 30 days in culture, Mock, Donor Only, or half of the GALC targeted cells were harvested for gDNA, while the other half of GALC targeted cells were CD19 enriched and expanded for another 7 days before gDNA harvest. N=3-5.  C) gDNA from WT fibroblasts or Krabbe Disease fibroblasts from two patients (purchased from Coriell; 6806 and 8304) were analyzed for GALC deletion as described in the materials and methods. Cell line 6806 is a homozygous knockout for the deletion and 8304 is heterozygous for the GALC deletion. D) GALC-K562 cells were generated to establish the GALC enzyme activity assay. These data show that the 6806 and 8304 Krabbe fibroblast cell lines are deficient in GALC enzyme activity. Data are represented as mean +/- SEM.
Supplemental Figure 7: GALC-NSCs engraft in the mouse cerebellum and produce myelin in vivo. A) immunoperoxidase staining with the human-specific mAb SC121 (brown with red arrow) detects engraftment of human cells in the white matter in the cerebellum. B) immunoperoxidase staining of a sibling section with anti-MBP (brown with red arrows) reveals a similar distribution of grafted GE-NSCs. Scale bars shown are 500µm. These data show that human cells that engrafted after 8 weeks are producing myelin.
Transparent Methods:

**NSC cell culture**

Human NSCs grown as neurospheres were generated as described under non-GMP conditions (Uchida et al., 2000, Uchida et al., 2012). were cultured at a density of 1E5 per ml in X-VIVO 15 medium (Lonza) supplemented with N2 (Invitrogen, 1:100), heparin (2 ug/ml), N-acetylcysteine (63ug/ml), fibroblast growth factor 2 (20 ng/ml), epidermal growth factor (20 ng/ml), and leukemia inhibitory factor (10 ng/ml). Neurospheres with Passage 8-13 were used in this study. Neuropsheres were disoociated by purified collagenase (Liberage/Blenzyme, Rohche) into single cell suspension and replated for passage in the medium described above.

**Genome Editing Reagents:**

**TALEN-mediated genome editing**: IL2RG TALENs were synthesized (GenScript) using the Δ152 N-terminal domain and the +63 C-terminal domain and fused to the FokI nuclease domain and cloned into pcDNA3.1 (Invitrogen) as described (Hendel et al., 2014). **CRISPR/Cas9-mediated genome editing**: sgRNA expression vectors were constructed by cloning of 20 bp oligonucleotide target sequences into px330-U6-Chimeric_BB-CBh-hSpCas9 (a kind gift from Feng Zhang, Addgene plasmid #42230) containing a human codon-optimized SpCas9 expression cassette and a human U6 promoter driving the expression of the chimeric sgRNA (Hendel et al., 2014). The genomic target sequences for the guides were as follows: HBB: 5′-CTTGCCCCACAGGCGCATGAA-3′, CCR5: 5′ GCAGCATAGTGAGCCAGAA-3′, IL2RG: 5′ -GGTAATGATGGCCTTCAACA-3′. Chemically modified sgRNAs: Cas9 was delivered as 5meC, Ψ-modified mRNA (TriLink BioTechnologies) and sgRNAs were synthesized by Agilent Technologies (MS and MSP), Synthego (MS) or TriLink BioTechnologies (MS) with 2′-O-methyl (M), 2′-O-methyl 3′phosphorothioate (MS), or 2′-O-methyl 3′thioPACE (MSP) incorporated at three terminal nucleotides at both the 5′ and 3′ ends, and thus represents the “All RNA” delivery of the CRISPR system (Hendel et al., 2015). **Targeting vectors**: CCR5 and IL2RG plasmid targeting vectors had ~2 × 800 bp arms of homology, which were generated by PCR amplification of the corresponding
loci using genomic DNA isolated from K562 cells. *HBB* had homology arms of 540bp and 420bp. The homology arms were cloned into a ~2,900 base pair vector based on pBluescript SK⁺ using standard cloning methods (Hendel et al., 2015, Hendel et al., 2014). Between the homology arms, donors contain the Ubiquitin C promoter (UbC) driving expression of GFP (for both HBB plasmid and AAV6 donors). For CCR5 biallelic editing, one plasmid construct had UbC-GFP and the other had UbC-mCherry-T2A-tCD19. The *IL2RG* homologous donor construct had an *IL2RG* cDNA upstream of the UbC promoter and GFP (See Figure 1A). Alternatively, bicistronic cassette constructs separating proteins by a self-cleaving 2A peptide were generated as follows: GFP-2A-CD8 (CD8 alpha cell surface marker for purification of GE-NSCs), GFP-2A-tCD19 (truncated cell surface marker for purification of GE-NSCs) and GALC-2A-tCD19 (therapeutic enzyme construct for Krabbe disease with tCD19).

**Genome Editing of NSCs**

Single suspension of 500,000 NSCs were transfected with 1 µg TALEN-encoding plasmids and 1-2 µg donor plasmid (unless otherwise indicated) by nucleofection (Lonza) with an Amaxa 4D Nucleofector (program CA137) with the P3 Primary Cell Nucleofector Kit (V4XP-3032) using 20 µL, 16-well Nucleocuvette strips following manufacturer’s instructions. Alternatively, 2.5 x 10⁶ NSCs were transfected with 5 µg TALEN-encoding plasmids and 5-10 µg donor plasmid (unless otherwise indicated) by nucleofection (Lonza) with an Amaxa 4D Nucleofector (program CA137) with the P3 Primary Cell Nucleofector Kit (V4XP-3024) using 100 µL Nucleocuvettes following manufacturer’s instructions. For experiments with chemically modified sgRNAs and Cas9 mRNA, 500,000 NSCs were electroporated with 15µg Cas9 mRNA and 10µg MS or MSP sgRNA. After nucleofection, NSCs were plated in culturing flasks and cultured for multiple passages.

**Genome Editing of NSCs with Cas9 mRNA and AAV6 donors**

A single cell suspension of 500,000 NSCs were responded with 15µg of Cas9 mRNA was mixed with 10µg of sgRNA in 20 uL of P3 solution (Lonza). Nucleofection was performed using 16-well Nucleocuvette Strip with the 4D Nucleofector system (Lonza).
using CA137 code. Immediately after electroporation, cells were transferred into one well of a 48 well plate containing 250 µl of NSC media. Then the HBB UbC-GFP AAV6 donor vector (purchased from Vigene Biosciences) was added directly to the electroporated cells at vector genomes/cell of 10,000, 100,000, or 500,000. After 24 hours, cells were transferred into a T25 flask with 5ml of NSC media and cells were harvested and gDNA was extracted 6 days later (7 days total post electroporation).

**Analysis of homologous recombination (HR) via flow cytometry and ‘In-Out’ PCR**

To monitor homologous recombination (HR), GE-NSCs were analyzed for GFP expression at each passage after nucleofection. GFP expression was measured on an Accuri C6 flow cytometer (BD Biosciences, San Jose, CA, USA). Alternatively, expression of transgene CD8 or CD19 was assessed by flow cytometry at each passage. At passage, neurospheres were dissociated as described above and single cell suspension were immunostained with anti-CD8-APC or anti-CD19-APC (Miltneyi Biotech) following manufacture’s instructions. ‘In-Out’ PCR was performed to qualitatively identify integration at the intended genomic locus where one primer binds outside the homology arms and the other primer binds inside the donor cassette; thus amplification should only occur if the donor cassette is integrated correctly.

**Digital Droplet PCR (ddPCR) analysis of on-target integration of HR donors**

ddPCR was carried out to determine the frequency of allelic targeted integration in human NSCs as previously described (Vakulskas et al., 2018). Briefly, genomic DNA was extracted using QuickExtract DNA Extraction solution (Epicenter) and then digested in either HINDIII-HF (for analyzing 3’ integrations into HBB) or BamH1-HF (for analyzing 3’ integrations into IL2RG or CCR5) for 2 hours. 3µl of digested gDNA was then used as the template for the ddPCR reaction. Droplet samples were prepared according to manufactures protocol (Bio-Rad) and PCR cyclin conditions were used as follows: 98°C (10 min); 94°C (0.5 min); 60°C (0.5 min); 72°C (2 min; 50 cycles); and 98°C (10 min). Finally, droplets were analyzed according to the manufacturer’s instructions using the QX200 system (Bio-Rad). The gene-specific integrations primer/probes were designed to analyze BGH PolyA signal at the 3’ end of the
recombinant alleles. The primers used to amplify the gene-specific targeted alleles and reference alleles were as follows: HBB forward: 5'-GGGAAGACAATAGCAGGCAT-3’, HBB reverse: 5’-CGATCCTGAGACTTCCACAC-3’, HBB probe: 5’-FAM-TGGGGATGCAGTGGGCTCTATGGC-BHQ-3’, CCR5 forward: 5’-GGGAAGACAATAGCAGGCAT-3’, CCR5 reverse: 5’-TCAAGAATCAGCAATTCTCAGGC-3’, CCR5 probe: 5’-FAM-TGGGGATGCAGTGGGCTCTATGGC-BHQ-3’, IL2RG forward: 5’-GGGAAGACAATAGCAGGCAT-3’, IL2RG reverse: 5’-CAGATATCCAGAGCCTAGCCTCATC-3’, IL2RG probe: 5’-FAM-TGGGGATGCAGTGGGCTCTATGGC-BHQ-3’, CCRL2 (reference) forward: 5’-GCTGTATGAATCCAGGTCC-3’, CCRL2 reverse: 5’-CCTCCTGAGCGTTGAGAAAAG-3’, CCRL2 probe 5’-HEX-TGTTTCTCCAGGATAAGGCAGCTGT-BHQ-3’.

INDEL analyses using tracking INDELs by decompression (TIDE) software
Genomic DNA (gDNA) was extracted at least seven days following electroporation of engineered nucleases in NSCs. Gene-specific primers were used to amplify alleles of interest (CCR5, HBB, IL2RG) and purified PCR fragments were then Sanger-sequenced as described(Dever et al., 2016). INDEL frequencies were quantified using the TIDE software (Tracking of Indels by Decomposition)(Brinkman et al., 2014).

Restriction Fragment Length Polymorphism (RFLP) Allele Analysis
The introduction of an AflII restriction site between 800bp homology arms allowed analysis of HR frequencies by Restriction Fragment Length Polymorphism (RFLP) at the IL2RG locus. Cas9 (under CMV promoter) + sgRNA (under U6 promoter) was delivered in the px330 plasmid construct (1ug). TALEN pairs were constructed as described above using the golden gate cloning system and delivered as plasmid constructs (0.5ug of each). 500,000 NSCs were nucleofected with either 1ug HR donor or 1ug HR donor and a nuclease. Cells were allowed to grow in culture for 7 days, gDNA was harvested, In-Out PCR was performed, the PCR product was isolated, and finally digested with AflIII overnight and products were run on 10% PAGE gels and visualized and quantified for band intensities via ImageJ software. The amount of modified IL2RG alleles was
quantified by dividing the density of cut alleles (800bp) by total alleles (unmodified (1.6kb) and modified).

**Purification of GE-NSCs by Magnetic Activated Cell Sorting (MACS)**

To Purify GE-NSC using either CD8 or CD19 transgene cell surface marker, either human CD8 Microbeads or CD19 MicroBeads (Milenyi Biotec) was used with following the manufacturer's instructions. Briefly, NSCs were expanded 4-6 passages after transfecting plasmids containing, Cas9, sgRNA and targeting vectors. Cells were harvested and dissociated to single cell suspensions by blenzyne and cell number was determined. Cells were centrifuged, resuspended into MACS buffer and added either CD8 or CD19 Microbeads according to manufacturer’s protocol. Magnetic separation was performed by using MS column to collect magnetically labeled cells. The eluted cells were applied to a second MS column to increase the purity of CD8+ or CD19+ cells. The purity of MACS selected cells were examined by staining either antibodies against CD8-APC or CD19-APC (Miltenyi Biotec). The purified cells were for further expanded and analyzed at each passage.

**Transplantation of GE-NSC**

Shi mice (C3Fe.SWV-Mbpshi/J) were backcrossed with immunodeficient Rag2/IL-2Rγ knockout mice (Shi-id). For transplantation, a suspension of GE-NSCs (1E5 cells in 1 microL per site) was prepared and transplanted bilaterally into the corpus collusum, SVZ and cerebellar white matter of neonatal or juvenile Shi-id mice as described previously(Uchida et al., 2012). A total of 114 mouse brains (80 females, 34 males) were analyzed for human cell engraftment by human specific monoclonal antibody SC121 (Fig 3A, 3E and Supplemental 7) (see staining details below). After evaluating human cell engraftment, 11 mouse brains were stained for mAb against GFP for transgene expression (Fig 3A and 3F). hNSC-derived myelin/myelinating oligodencytes were tested in 34 homozygous shiverer mouse brains by staining antibody against human MBP, which did not stain for mouse MBP in shiverer brains (Fig 3G and 3H). Three mouse brains were analyzed for Sox2, GFAP and DCX along with GFP (Fig 3B-D). There were no differences observed between females and males for human cell engraftment and
transgene expression. All animal housing conditions and surgical procedures were approved by and conducted according to the Institutional Animal Care and Use Committee at StemCells, Inc.

**Histology**

Transplanted mice were anesthetized and perfused with PBS followed by 4% paraformaldehyde. Brains were serially sectioned (50µm) in the sagittal plane with a freezing microtome (Leica SM2400). For immunoperoxidase staining with SC121 (1:1000, StemCells Inc.), anti-MBP (Millipore 1:500), and anti-GFP (1:500, Invitrogen), brain sections were stained with primary antibodies, followed by biotinylated horse anti-mouse mAb secondary antibodies (1:500, Vector Laboratory). Peroxidase staining was developed with an Elite ABC kit (Vector Laboratory) with NovaRed substrate (Vector Laboratory). Brain sections were mounted and counterstained with methyl green. For confocal immunofluorescence microscopy, imaging was performed on a Leica SP2 AOBS microscope (Leica Microsystems) or a LSM 780 (Zeiss). Brain sections were stained with antibodies against GFP, MBP, Sox 2 (R&D System, 1:200), SC123 (human GFAP, StemCells. Inc 1:3000), and Doublecortin (Santa Cruz 1:200). Secondary antibodies were incubated at RT for 2 hours: Alexa Fluor 488 donkey anti-rabbit (1:500 Invitrogen), Alexa Fluor donkey anti-goat 568 (1:500 Invitrogen), Alexa donkey anti-mouse 568 (1:500, Invitrogen) and counterstained with Hoechst or DAPI to identify nuclei.

**Targeted amplicon library generation for MiSeq runs**

*IL2RG* (ON) and the top three *in silico* predicted off target (OFF1-3) Cas9-sgRNA amplicons were PCR-amplified with sequencing primers utilized in deep sequencing MiSeq runs as previously reported(Hendel et al., 2015). Amplicons were gel purified and then subjected to a second round of PCR to add adapters and unique 8bp barcodes to distinguish experiments. Barcoded amplicons were then purified and pooled in equimolar concentrations. The purified library was sequenced on an illumina MiSeq DNA sequencer at 2 x 200 cycles with indexing at the Protein and Nucleic Acid (PAN)
Stanford Core Facility. Sequences were aligned to the human genome and INDELs were calculated as described below.

**MiSeq analyses of in silico predicted IL2RG sgRNA off-target sites**

*IL2RG* (ON) and top three *in silico* predicted OFF-target Cas9-sgRNA sites were quantified by mapping reads from each sample to the four amplicon sequence target regions (*IL2RG*, OFF1-3) and measuring number of mapped reads with a gap in a 10 base pair neighborhood of the cut site. Specifically, each read was first assigned to one of the target regions by finding a perfect match between the first 70 bases of each read and the amplicon sequence. If the first 70 base pairs of a read do not perfectly map to any amplicon sequence the read is discarded, resulting in the removal of a large number of low-quality reads from each sample. Reads that do not span a whole amplicon are also discarded. Each read was then aligned with its corresponding target amplicon sequence using EMBOSS version 6.5.7.0 needle with default parameters (Rice et al., 2000). To quantify the number of reads with INDELs, each read was marked as modified if at least one insertion or deletion (a “-“ in the alignment of either read or the amplicon) occurs within 5bp up or downstream of the CRISPR cut site (between bases 17 and 18 of the guide RNA sequence). The overall INDEL percentage at a given target site was reported as the number of modified reads mapping to the locus over the total number of reads mapping to the locus, minus the background INDEL percentage in a non-electroporated sample.

**GALC enzyme cross-correction of GALC-deficient Krabbe disease Fibroblasts**

Fibroblasts from two patients with infantile Krabbe disease were purchased from Coriell Institute for Medical Research (GM06806 and GM08304). Both Krabbe disease fibroblast cell lines have a large deletion in *GALC* that results in deficient GALC enzyme activity (*Supplemental Figure 6C and 6D*), which has been previously reported (Rafi et al., 1995, Luzi et al., 1995). To assess whether GALC-NSCs could rescue GALC activity in Krabbe disease Fibroblasts, we performed cross-correction experiments where GALC-NSCs were co-cultured in 3µm transwell permeable polyester membrane inserts (Corning) with Krabbe fibroblasts. To achieve this, 100,000 fibroblasts were seeded
overnight in NSC media and allowed to adhere and on the following day 1,000,000 GALC-NSCs were added to the permeable membrane insert. Cells were co-cultured for 7 days and half of the fibroblasts were harvested for GALC enzyme activity (passage 1) and the other half were seeded overnight in NSC where on the following day fresh GALC-NSCs were co-cultured in the membrane insert. After 7 more days of cross-correction (and 14 days total), Krabbe fibroblasts were harvested, protein was isolated and GALC enzyme activity was performed as described below. To inhibit GALC enzyme uptake, mannose-6-phosphate (Sigma) was added to the media every 24-48h at a final concentration of 2.5mM.

**GALC enzyme assay**

Cellular protein was extracted by lysing cells in deionized water with a Branson Sonicator with probe, centrifuging lysates at 17,000xg for 10 minutes at 4°C, and collecting the supernatant containing the soluble proteins. Protein concentration in the supernatants was measured by Bradford assay kit with BSA standard curve ranging from 0.25-5 mg/ml (Thermo Scientific). To prepare the GALC assay substrate, 250 uL of .9mM 6-hexadecanoyl-4-methylumbelliferyl-beta-D-galactopyranoside (Sigma) in chloroform/methanol (2/1 v/v), 75 uL of 6mg/mL oleic acid in hexane, and 50 uL of 30mg/mL sodium taurocholate in chloroform/methanol (2/1 v/v) were combined and the mixture was dried using a rotary evaporator. The dried mixture was stored under nitrogen at -80 °C covered from light until ready for use. The working substrate was prepared by resuspending the dried substrate in 500 uL assay buffer (50 mM citrate, 100 mM phosphate, pH 5.0) to a final concentration of 0.45 mM and sonicating briefly. 25-50 ug protein extract (50 uL) was mixed with 100 uL of working substrate and incubated for 20 hours at 37 °C covered from light. Reactions were stopped with 200 uL stop buffer (0.2 M glycine/NaOH, 0.2% w/v sodiumdodecyl sulfate, 0.2% w/v Triton X-100, pH 10.7). Fluorescence of 4-methylumbeliferone (4MU) liberated by GALC enzyme cleavage was measured using a Molecular Devices SpectraMax M3 multi-mode microplate reader with SoftMax Pro 5 software at excitation and emission wavelengths of 355 nm and 460 nm,
respectively (top read). A standard curve for 4MU was established using 4MU sodium salt (Sigma) in assay buffer.

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