Elements of lentiviral vector design toward gene therapy for treating mucopolysaccharidosis I

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Abstract

Mucopolysaccharidosis type I (MPS I) is a lysosomal disease caused by α-L-iduronidase (IDUA) deficiency and accumulation of glycosaminoglycans (GAG). Lentiviral vector encoding correct IDUA cDNA could be used for treating MPS I. To optimize the lentiviral vector design, 9 constructs were designed by combinations of various promoters, enhancers, and codon optimization. After in vitro transfection into 293FT cells, 5 constructs achieved the highest IDUA activities (5613 to 7358 nmol/h/mg protein). These 5 candidate vectors were then tested by injection (1 × 10⁷ TU/g) into neonatal MPS I mice. After 30 days, one vector, CCEoIDW, achieved the highest IDUA levels: 2.6% of wildtype levels in the brain, 9.5% in the heart, 200% in the liver and 257% in the spleen. CCEoIDW achieved the most significant GAG reduction: down 49% in the brain, 98% in the heart, 100% in the liver and 95% in the spleen. Further, CCEoIDW had the lowest transgene frequency, especially in the gonads (0.03 ± 0.01 copies/100 cells), reducing the risk of insertional mutagenesis and germ-line transmission. Therefore, CCEoIDW is selected as the optimal lentiviral vector for treating MPS I disease and will be applied in large animal preclinical studies. Further, taken both in vitro and in vivo comparisons together, codon optimization, use of EF-1α promoter and woodchuck hepatitis virus posttranscriptional response element (WPRE) could enhance transgene expression. These results provided a better understanding of factors contributing efficient transgene expression in lentiviral gene therapies.

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1. Introduction

Mucopolysaccharidosis type I (MPS I) is an autosomal recessive disease that leads to neurodegeneration, mental retardation and death in early age. MPS I results from deficiency of α-L-iduronidase (IDUA, EC 3.2.1.76), which degrades the glycosaminoglycans (GAG). The pathological study of MPS I is extremely difficult to treat, because the blood-brain-barrier (BBB) blocks the entry of enzyme into the brain [1–3]. Currently, MPS I is treated by hematopoietic stem cell transplantation (HSCT) and often in conjunction with enzyme replacement therapy (ERT). However, HSCT has a high rate of mortality (10–15%) and severe morbidity. While ERT alone is thought to have negligible therapeutic impacts at the brains, a high-dose infusion of IDUA can lead to significant increase of enzyme activity in the brain cortex [4]. Given that a small amount of enzyme is sufficient to degrade GAG storage [5–7], it should be possible to achieve improved neurological outcomes. However, high dose treatment is not feasible for clinical application due to the immune response against human recombinant enzyme [8].

Previously, gene therapy with different vectors in animal models has been used to treat MPS I disease, including retrovirus [9,10], lentivirus [8,11,12], adeno-associated virus [3,13–15], Sleeping Beauty transposon [16] and minicircles [17]. It has been shown that injection of lentiviral vector into MPS I mice can achieve metabolic correction and neurological improvements [8,11,12]. However, it is difficult to apply the dose (1.65 to 4.5 × 10⁸ TU/g) used into clinical trials. Therefore, optimization of lentiviral vector design is essential for advancing lentiviral gene therapy protocols.

Promoters and enhancers are essential vector components for optimizing transgene expression. It has been shown that the human phosphoglycerate kinase 1 (PGK) promoter leads to moderate transgene expression of lentiviral vector [18–20]. Lentiviral vector with the human elongation factor 1α (EF-1α) promoter has been used for treating Fabry disease in a murine model [21]. Additionally, a hybrid promoter consisting of the enhancer of the murine cytomegalovirus (CMV) immediate-early gene and human EF-1α promoter was shown to achieve high transgene expression of lentiviral vector [22]. However, the effects of different promoters on transgene expression are still not elucidated. Moreover, the use of woodchuck hepatitis virus (WHV) posttranscriptional response element (WPRE) has been found to enhance transgene expression and titers of therapeutic vectors [23–25]. The enhancing ability of WPRE depends on target cells, the type of viral vector context and its sequence [26–29]. However, WHV X protein is implicated in the development of liver tumors [30], which raises the safety concern about use of WPRE in vectors for gene therapy. Herein,
we designed constructs with full-length WPRE, truncated WPRE (tWPRE) and depleted WPRE for a side-by-side comparison. In this study, an initial in vitro screening of 9 plasmids identified 5 candidates with the highest IDUA transgene expression. Then, the efficacy of these 5 candidate lentiviral vectors in neonatal MPS I mice was comparatively evaluated. This allowed us to determine which lentiviral constructs yielded the highest IDUA levels and the most efficient GAG reduction in vivo.

2. Materials & methods

2.1. Plasmid construction

The human IDUA cDNA generated by reverse transcription PCR from total mRNA of an unaffected individual was inserted into the multi-cloning sites of pHIV-CS (CMV promoter upstream of 5′ LTR). The IDUA expression was under the control of PGK promoter and named as pCPGKID. Then, codon optimization of IDUA cDNA sequence was performed, resulting in what we named the oIDUA sequence. Similar techniques were used to generate similar variants with different promoters (hybrid promoter named as CE, PGK and EF-1α) and variants of WPRE, resulting in 8 more plasmids as followed: pCEFDW, pCEFIDW, pCPGKDw, pC EvoDW, pCEF0ID-wPRE, pCEFID, pCEID and pCEIDW. All plasmids were confirmed by sequencing.

2.2. In vitro transfection

For each transfection, 25 μg of candidate plasmid and 25 μg of HIV CMVeGFP plasmid were mixed with 133 μL 2.5 M CaCl₂ (25 °C) and 1.33 ml RNase/DNase free sterile H₂O. After adding 1.33 ml of 2× HEPES buffered saline (pH 7.1), 7 ml serum free medium was added to the mixture. Then, the HEK 293FT cells were incubated with this transfection solution for 4 to 6 h (37 °C, 5% CO₂). After removing the transfection solution, cells were incubated with 9 ml of 10% fetal bovine serum (FBS) (Sigma-Aldrich, St. Louis, MO), Dulbecco's modified eagle medium (DMEM) (Sigma-Aldrich, St. Louis, MO) was used to generate the standard curve. The resulting fluorescence was measured with excitation at 355 nm and emission at 460 nm. IDUA enzyme activity was expressed in units (nmol converted to product per hour) per mg protein as determined with a Pierce protein assay kit (Fisher, Waltham, MA) or per ml plasma. Then, IDUA enzyme activity was adjusted by Michaelis–Menten equation as described previously [33]. All reactions were performed in triplicate.

2.3. Lentiviral vector production

The candidate plasmid was co-transfected with three additional helper plasmids, plP1 (gag/pol), plP2 (Rev) and VSVG envelope plasmid into HEK 293FT cells [31]. Vector-containing medium was collected into HEK 293FT cells [31]. Vector-containing medium was collected by centrifuge and processed for biochemical assays.

2.4. MPS I mice and injection

MPS I knockout mice (idua−/−), a kind gift from Dr. Elizabeth Neufeld, UCLA, were generated by insertion of neomycin resistance gene into exon 6 of the 14-exon IDUA gene on the C57BL/6 background [32]. The colony was housed in a pathogen-free facility on a 12-hour light/dark cycle. Newborn mice (1–2 days old) were injected with concentrated lentiviral vector through the superficial temporal vein. All mouse care and handling procedures were in compliance with the rules of the Institutional Animal Care and Use Committee (IACUC) of the University of Minnesota.

2.5. IDUA enzyme assay

IDUA enzyme assay was conducted as previously described [33]. IDUA activity was determined by a fluorometric assay using 4-methylumbelliferyl α-L-iduronide (4-MU iduronide) (Glycosynth, Cheshire, UK) as the substrate, which was diluted with sodium formate buffer (0.4 M, pH 3.5). Then, 25 μl aliquots (360 μM) of substrate were mixed with 25 μl aliquots of tissue homogenates (diluted with 0.2% bovine serum albumin in phosphate buffered saline). The mixture was incubated at 37 °C for 30 min, and 200 μl glycerine carbonate buffer (pH 10.4) was added to stop the reaction. IDUA catalyzed the cleavage of the non-fluorescent substrate (4-MU iduronide) and released a fluorescent product (4-MU). 4-Methylumbelliferone (4-MU) (Sigma-Aldrich, St. Louis, MO) was used to generate the standard curve. The resulting fluorescence was measured with excitation at 355 nm and emission at 460 nm. IDUA enzyme activity was expressed in units (nmol converted to product per hour) per mg protein as determined with a Pierce protein assay kit (Fisher, Waltham, MA) or per ml plasma. Then, IDUA enzyme activity was adjusted by Michaelis–Menten equation as described previously [33]. All reactions were performed in triplicate.

2.6. Tissue GAG assay

Tissue GAG assays were conducted as described previously [4]. The supernatants of tissue homogenates were treated by proteinase K (ProK) (NEB, Ipswich, MA) with the ratio of 3 (Pro K):1 (sample), incubated at 55 °C overnight, and boiled for 10 min to inactivate the enzyme. Samples were incubated with 2.5 μg RNase (Sigma-Aldrich, St. Louis, MO) and 250 U DNase (Sigma-Aldrich, St. Louis, MO) at room temperature overnight. After boiling for 10 min to inactivate the enzymes, GAG levels were determined by the Blyscan Sulfated Glycosaminoglycan Assay (Biocolor, Carrickfergus, UK). Tissue GAG levels were expressed as μg GAG/mg protein.

2.7. Quantitative PCR

DNA was extracted from tissues with QIAGEN DNA mini kit (QIAGEN # 51306). Primers (forward primer: 5′-CGACTGGTGAGTACGCCAAA-3′; reverse primer: 5′-CCGACCATCTCCTCCTCT-3′) and the probe (5′-FAM-ATTTTGACTAGCGGAGG-TAMRA-3′) targeted the lentiviral psi (Ψ) packaging signal region. Amplicon size was 61 bp. Each reaction contained 2 × TaqMan universal PCR master mix (Life Technologies, Carlsbad, CA), primers (0.008 nmol each), TaqMan probe (0.008 nmol), and 100 ng of sample DNA in a final volume of 10 μL. Real-time PCR was also performed using the mouse apolipoprotein B (ApoB) gene as an internal control, using the following primers (forward primer, 5′-CGTGGGCTCCAGATCCTC-3′; reverse primer, 5′-TCACCGT CATTCTGGCTTTT-3′) and probe (5′-FAM-CCTTGACAGGTGCCCAACCA TTC-TAMRA-3′). Amplification conditions were 2 min at 50 °C and 10 min at 95 °C for the first cycle, followed by 50 cycles of 95 °C for 15 s and 60 s for 1 min. A standard curve was established from a series of genomic DNA mixtures derived from plasmid with Ψ packaging signal region sequence (1 copy per genome). Unknown samples were run in triplicate, while standards and internal controls were performed in duplicate.

2.8. Statistical analysis

Data were represented as mean ± standard errors. For evaluation of differences between samples, Student’s T test for comparisons between paired samples and one-way analysis of variance (ANOVA) for comparisons between three or more samples were performed. Statistical significant level was set at p < 0.05. Data analysis was conducted with SAS 9.3 (Cary, NC).
3. Results

3.1. Design of 9 lentiviral constructs

A plasmid CPGKID was constructed using the human PGK promoter to drive expression of human IDUA cDNA in the genome of a self-inactivating (SIN) lentiviral vector. Through codon optimization, optimized IDUA (oIDUA) sequence was obtained and used in 5 constructs. EF-1α promoter and the hybrid promoter containing CMV enhancer and EF-1α were engineered into 4 and 3 different constructs, respectively. Full length of WPRE, which contains three sub-elements: α, β and γ, was introduced into 5 different constructs. It is known that sub-elements γ and α contribute to most of the enhancing function, while β contains WHV X protein [34]. Due to safety concern about the WHV X sequence, CEFoID-tWPRE is designed by deleting β sub-element. The sequence information of these 9 lentiviral constructs was represented in Fig. 1.

3.2. Comparison by in vitro transfection

Each of these 9 plasmids was transfected into HEK 293FT cells, and three independent transfection experiments were conducted. The transfection efficiency of each plasmid was similar, shown by co-transfection of a plasmid encoding GFP (data not shown). A total of 5 constructs (CEFoIDW, CEFoID-tWPRE, CEFoID, CCEoIDW and CPGKID) yielded the highest IDUA levels in cell lysates (Fig. 2). Also, these 5 constructs had the highest IDUA levels in supernatants, which confirmed the results seen in the cell lysates (Fig. 2). These 5 constructs were

Fig. 1. Graphical representation of 9 lentiviral vectors. The lentiviral backbone is derived from HIV-1 with self-inactivating (SIN) LTRs. Rev. response element (RRE), central polypurine tract (cPPT) and central termination sequence (CTS) are in all constructs but not shown here. LTRs: long terminal repeats; Ψ: HIV-1 packaging signal; oIDUA stands for codon optimized version of human IDUA cDNA sequence; PGK: human phosphoglycerate kinase 1 promoter; EF-1α: human elongation factor 1α promoter; CE: mouse CMV enhancer and human EF-1α promoter; IDUA: human IDUA cDNA; oIDUA: codon optimized human IDUA cDNA; WPRE: WHV posttranscriptional response element; tWPRE: truncated WPRE by deleting β sub-element.

Fig. 2. IDUA enzyme levels in cell lysates and supernatants 48 h after transfection of candidate plasmids. IDUA enzyme activity in HEK 293 FT cells without transfection is <0.01 nmol/h/mL in supernatants, and 0.97 ± 0.34 nmol/h/mg protein in cell lysates, while IDUA enzyme activity in cultured medium without cells is 0. Data are mean ± standard errors (n = 3).
selected as candidates and packaged into lentiviral vectors for in vivo assessment. When comparing constructs with or without codon optimization, CEFoIDW yielded higher IDUA levels than CEFIDW (7359 ± 956 vs 4879 ± 947 nmol/h/mg protein, p < 0.05), while CCEoIDW achieved higher IDUA levels than CCEIDW (7334 ± 858 vs 3784 ± 656 nmol/h/mg protein, p < 0.05). As to comparison between constructs with different promoters, CEFoIDW and CCEoIDW had IDUA levels at 7359 ± 956 and 7334 ± 858 nmol/h/mg protein, respectively, which is higher than that of CPGKoIDW (3732 ± 106 nmol/h/mg, p < 0.05). To determine the effects of WPRE, side-by-side comparisons (CEFoIDW vs CEFoID-tWPRE vs CEFoID) were conducted, and CEFoID had slightly lower IDUA levels than the other constructs (not statistically significant). This observation was confirmed when comparing CCEIDW (3732 ± 106 nmol/h/mg protein) and CCEoID (3221 ± 302 nmol/h/mg protein). Notably, the ratio between IDUA levels in supernatants and cell lysates was approximately 11% in all 9 plasmids, indicating that the overexpression of IDUA did not affect the normal secretion of IDUA.

A previous study with retroviral gene therapy observed the inclusion-cell (I-cell) phenotype when IDUA levels were extremely high within cells [35]. It was proposed that due to saturation of M6P mediated lysosome-targeting pathway by overexpressed IDUA, other lysosomal enzymes could not be efficiently delivered to the lysosome. To test this possibility, enzyme activity of iduronate-2-sulfatase (IDS) in transfected cells and supernatants were assessed. With high IDUA levels, IDS levels in cell lysates decreased to between 2.5 and 3.5 nmol/h/mg protein (wildtype level: 5.5 nmol/h/mg protein), and IDS levels in supernatants increased to approximately 1 nmol/h/mL (wildtype level: 0.89 nmol/h/mL). These results showed that targeting of IDS to the lysosome is only slightly affected by overexpressed IDUA.

### Table 1
Quantitative measurement of lentiviral DNA in tissues of MPS I mice with 5 candidate lentiviral vectors. The untreated MPS I mice had 0.014 copies/100 cells in the liver, non-deetectable (ND, < 0.004%) to 0.023 in the spleen, ND in the brain, and ND to 0.006 in the gonad. Data are mean ± standard errors.

| Copies/100 cells | Brain | Gonad | Liver | Spleen |
|------------------|-------|-------|-------|--------|
| CEFoIDW (n = 5)  | 0.16 ± 0.06 | 0.15 ± 0.1 | 6.4 ± 0.42 | 2.4 ± 0.78 |
| CEFoID-tWPRE (n = 7) | 0.24 ± 0.09 | 0.1 ± 0.04 | 9.8 ± 3.3 | 3.7 ± 1 |
| CCEoIDW (n = 4)  | 0.06 ± 0.02 | 0.03 ± 0.01 | 4.9 ± 2.5 | 1.5 ± 0.76 |
| CPGKID (n = 4)   | 0.25 ± 0.16 | 0.01 ± 0.01 | 2.6 ± 0.04 | 2 ± 0.77 |
| CEFoID (n = 3)   | 0.43 ± 0.16 | 0.31 ± 0.16 | 5.7 ± 1.9 | 6.2 ± 1.7 |

3.3. In vivo comparison in neonatal MPS I mice

Vector particles were produced by co-transfection in HEK 293FT cells using a third-generation (4-plasmid) packaging system. Although a previous study [25] found that WPRE can enhance titers of retroviral vector, no significant difference in titers between lentiviral vectors with or without WPRE was identified in our three independent batches of virus production (data not shown). The 5 candidate lentiviral vectors were separately injected into newborn MPS I litters through the temporal facial vein at a dose of 1 × 10⁷ TU/g (injection volume: 40 to 50 μL). Each group enrolled a litter of MPS I pups from MPS I parents (n = 4 to 7).

All mice including untreated MPS I controls (n = 5) were sacrificed at the age of 30 days. To determine the biodistribution of lentiviral vectors, genomic DNA from the brain, gonad, liver and spleen were extracted and assessed by quantitative PCR (QPCR) (see details in Table 1). The highest levels of IDUA were observed in the liver ranging from 2.6 to 9.8% (i.e., 2.6 to 9.8 IDUA copies per 100 cells), with 1.5 to 6.2% in the spleen. Of the tissues surveyed, the lowest transgene levels were observed in the gonads, ranging from 0.03 to 0.31%. Of all 5 candidate vectors, CCEoIDW achieved the lowest transgene frequency in the...
brain (0.06%), gonad (0.03%) and spleen (1.5%), indicating less risk for insertional mutagenesis and germ-line transmission. All 5 candidate vectors achieved significant IDUA enzyme activity in the liver (0.54 ± 0.07 to 7.2 ± 2.1 nmol/h/mg protein) and spleen (0.9 ± 0.18 to 52.5 ± 18.5 nmol/h/mg protein) (data summarized in Fig. 3). CEFoIDW and CCEoIDW achieved significant IDUA activity (0.15 ± 0.03 to 0.75 ± 0.36 nmol/h/mg protein) in the heart. Only CCEoIDW achieved higher IDUA activity in the brain (0.14 ± 0.07 nmol/h/mg protein, p = 0.053). CCEoIDW had the highest IDUA enzyme activity in all tested tissues. Notably, no correlation between transgene frequency and IDUA levels were observed. All 5 candidate vectors achieved significant GAG reduction in the liver (by 88% to 100%, p < 0.05) (summarized in Fig. 4). Of the 5 vectors, all except CEFoID achieved significant GAG reduction in the heart (by 59% to 98%, p < 0.05). CEFoIDW and CCEoIDW achieved significant GAG reduction in the spleen (by 88% to 95%, p < 0.05). CEFoIDW and CCEoIDW achieved significant GAG reduction in the brain (by 34% to 38%, p < 0.05). Interestingly, although no significant IDUA levels were observed in the brain, CEFoID+tWPRE achieved significant GAG reduction (by 36%, p < 0.05). Of 5 candidate vectors, only CCEoIDW and CEFoIDW achieved significant GAG reduction in all tested organs, and CCEoIDW has the lowest GAG levels in the liver, spleen and heart. Taken together, with the lowest transgene frequency, the highest IDUA levels and the most significant GAG reduction, CCEoIDW emerges as the optimal lentiviral vector for treating MPS I disease.

### 3.4. Effects of dose and injection volume

To determine the effects of dose on treatment efficacy, a single dose of CPGKID (5 × 10^7 TU/g) was injected into neonatal MPS I mice (n = 8). The brain, heart, liver and spleen were collected for biochemical assays 30 days after the injection. Compared with a high dose of CPGKID (1 × 10^7 TU/g), the low dose group led to significantly lower IDUA levels in the liver (0.25 ± 0.05 vs 2 ± 0.54 nmol/h/mg protein, p < 0.05) and spleen (0.41 ± 0.08 vs 9.1 ± 3.7 nmol/h/mg protein, p < 0.05) (Table 2). There was no significant difference in GAG levels in the liver and spleen between these two groups. However, the low dose group led to significantly higher GAG levels (27.1 ± 2.9 vs 7.4 ± 0.84 μg GAG/mg protein) (Table 2). These results showed that the higher dose (1 × 10^7 TU/g) achieved more significant IDUA transgene expression and GAG reduction.

Previously, one study showed that when injecting gene vector into neonatal mice, the injection volume significantly affects transgene expression [36]. To determine the effects of injection volume on transgene expression, a single dose of CPGKID (1 × 10^7 TU/g) with the volume of

### Table 2

| IDUA activity (nmol/h/mg protein) | Volume (μL) | Brain | Heart | Liver | Spleen |
|---------------------------------|------------|-------|-------|-------|--------|
| CPGKID (high dose, n = 4)       | 50         | ND    | 0.09 ± 0.07 | 2 ± 0.54 | 9.1 ± 3.7 |
| CPGKID (low dose, n = 8)        | 50         | 0.006 ± 0.01 | 0.01 ± 0.01 | 0.25 ± 0.05 | 0.41 ± 0.1 |
| CPGKID (high dose, n = 5)       | 100        | ND    | 0.34 ± 0.13 | 1.7 ± 0.36 | 2.1 ± 0.07 |
| Untreated MPS I (n = 5)         | N/A        | ND    | ND    | ND    | ND     |

Data are mean ± standard errors. *The difference between treated mice and untreated MPS I mice was significant, with a p value of < 0.05.
heart, 7.4 ± 3.4 in the liver and 3.6 ± 1.0 in the spleen. Mice, both high volume and low volume groups achieved significant IDUA levels in all tested tissues (Table 2). Compared with untreated MPS I mice, there was no significant difference between these two groups, there was no significant difference in IDUA levels in all tested tissues (Table 2). Compared with untreated MPS I mice, both high volume and low volume groups achieved significant GAG reduction in the heart and liver (p < 0.05). However, there was no significant difference in GAG levels between these two groups (Table 3). Collectively, the higher injection volume did not achieve more significant IDUA levels and GAG reduction.

4. Discussion

4.1. Inference for lentiviral design

In this study, constructs with codon optimization achieved higher IDUA levels in vitro than their counterparts without codon optimization. Since there is no in vivo pairwise comparison, codon optimization can at least enhance transgene expression in vitro. In terms of promoters, EF-1α and the hybrid promoter achieved significantly higher IDUA levels than PGK in vitro. CCEoIDW achieved significantly the highest IDUA levels and efficient GAG reduction in vivo. There was no direct in vivo comparison between constructs with difference only in promoter choices (PGK or EF-1α). Taken together, the hybrid promoter containing CMV enhancer and EF-1α promoter performs better than PGK promoter both in vivo and in vitro. We also compared constructs with variants of WPRE (CEFoIDW vs CEFoID-tWPRE vs CEFoID, CCEIDW vs CCEID), there were no significant effects in vitro. In in vivo experiments, CEFoIDW achieved higher IDUA levels in the heart and more efficient GAG reduction than CEFoID. Similarly, when comparing with constructs with CEFoID-tWPRE, CEFoID had less efficient GAG reduction in the brain, heart and liver (p < 0.05). These results showed that although there were no significant effects in in vitro conditions, WPRE enhanced transgene expression and boosted GAG reduction in vivo.

4.2. IDUA activity and GAG reduction in the brain

Although the transgene frequency of all 5 lentiviral vectors in the brain is extremely low (0.06 to 0.43 copies/100 cells), some lentiviral vectors showed significant GAG reduction in the brain. This observation further confirmed the hypothesis derived from our high dose ERT study [4]: when plasma IDUA level is high enough, a small amount of IDUA can cross the blood–brain-barrier and degrade GAG storage in the brain. Another interesting observation is that CEFoID-tWPRE had undetectable IDUA levels in the brain, but achieved significant GAG reduction. Therefore, it appears that even a non-detectable amount of IDUA could significantly degrade GAG storage. This is consistent with previous findings in pseudo-deficient patients who have extremely low, undetectable IDUA levels, but with normal phenotype.

4.3. Clinical applications

Previous studies using lentiviral vector to treat MPS I disease is of limited of use due to relatively low transgene expression, making it difficult for translation into human clinical trials. Two studies [8,11] intravenously injected up to approximately 4.5 × 10^8 TU/g of lentiviral vector into adult MPS I mice. The liver had the most significant effects: only 1% of wildtype levels and 69% of GAG reduction. There were no IDUA activity nor GAG reduction in the brain. Another lentiviral gene therapy study [12] with neonatal MPS I mice used the dose of 1.65 × 10^8 TU/g, and achieved 27.4% of wildtype IDUA levels in the liver. However, for a 3 kg human baby, this dose means infusing approximately 2.5 L of lentiviral vector (set the vector titer as 2 × 10^9 TU/mL). In our study, we used the dose of 1 × 10^7 TU/g, namely 150 mL for human babies, which remarkably improved the ease of viral vector production. Moreover, CCEoIDW achieved more significant IDUA activity and efficient GAG reduction in all tested tissues including the brain.

In this study, with all 5 lentiviral vectors, the gonads always had the lowest transgene frequency (as low as 0.03 copies/100 cells) of all tested tissues. Similar to what a previous study showed [37], these results also indicated that the gonads are relatively isolated from intravenously administered vector, which remarkably reduced the risk of germ-line transmission. Another major safety concern about lentiviral vector is the oncogenesis by insertional activation. In this study, out of all tested tissues with all candidate vectors, the highest transgene frequency is 9.8 copies/100 cells 1 month after treatment (CEFoID-tWPRE in the liver). It is significantly lower than that in another lentiviral gene therapy study at the same time-point, which is 83 copies/100 cells [11]. Moreover, the optimal vector, CCEoIDW had the lowest transgene frequency: 0.06 in the brain, 0.03 in the gonad, 4.9 in the liver and 1.5 in the spleen. This makes CCEoIDW more appealing for significantly reducing risk of insertional mutagenesis. The optimal construct CCEoIDW has both CMV enhancer and WPRE, which may activate adjacent oncogenes and pose risks for insertional mutagenesis. In spite of these possibilities, these enhancer elements have been widely used in gene therapy without being linked to tumorigenesis.

Further, for future clinical trials, different routes of administration including in utero, intravenous, intranasal and intraventricular should be comparatively evaluated.

5. Conclusions

In conclusion, of 9 constructs, CCEoIDW emerges as the optimal lentiviral vector for treating MPS I mice and can be forwarded into preclinical studies. At the dose of 1 × 10^7 TU/g, CCEoIDW achieved supranormal IDUA levels and significant GAG reduction in tissues, and importantly in the brain. Considering the relatively low transgene frequency, this dose may be applicable to human clinical trials for neonatal injection. Further, inferences derived from this study about lentiviral vector design will be valuable for gene therapy studies treating other lysosomal diseases.
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