Full-length Dysferlin Transfer by the Hyperactive Sleeping Beauty Transposase Restores Dysferlin-deficient Muscle

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Dysferlin-deficient muscular dystrophy is a progressive disease characterized by muscle weakness and wasting for which there is no treatment. It is caused by mutations in DYSF, a large, multielexonic gene that forms a coding sequence of 6.2 kb. Sleeping Beauty (SB) transposon is a nonviral gene transfer vector, already used in clinical trials. The hyperactive SB system consists of a transposon DNA sequence and a transposase protein, SB100X, that can integrate DNA over 10 kb into the target genome. We constructed an SB transposon-based vector to deliver full-length human DYSF cDNA into dysferlin-deficient H2K A/J myoblasts. We demonstrate proper dysferlin expression as well as highly efficient engraftment (>1,100 donor-derived fibers) of the engineered myoblasts in the skeletal muscle of dysferlin- and immunodeficient B6.Cg-Dysf<sup>emd</sup> Prkdc<sup>Scid</sup>/J (Scid/BLA/J) mice. Nonviral gene delivery of full-length human dysferlin into muscle cells, along with a successful and efficient transplantation into skeletal muscle are important advances towards successful gene therapy of dysferlin-deficient muscular dystrophy.

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Subject Category: Gene insertion deletion or modification

Introduction

Miyoshi myopathy, limb girdle muscular dystrophy 2B, and distal anterior compartment myopathy are autosomal recessive muscle disorders caused by mutations in the gene encoding dysferlin (DYSF).<sup>1–3</sup> They are encompassed by the term dysferlinopathy and affect the proximal and/or distal muscles of the limbs. They generally appear after puberty and their clinical presentation is variable regarding the types of muscles involved and the degree of severity.<sup>1–3</sup> Patients usually become wheelchair-bound within 10–20 years of disease onset due to progressive muscle degeneration, weakness, and atrophy.

DYSF spans over 150 kb of genomic DNA in chromosome 2p13 and comprises 55 exons that form a coding sequence of 6.2 kb.<sup>1,2,4</sup> The 237 kDa dysferlin protein belongs to the ferlin family, a group of large proteins with important roles in vesicle trafficking and fusion.<sup>5</sup> DYSF is expressed in several tissues, including kidney and immune cells, but its highest expression was reported in muscle,<sup>6</sup> where dysferlin is mainly detected in mature myofibers.<sup>7</sup> In muscle fibers, dysferlin localizes predominantly to the sarcolemma, but it is also present at the transverse tubules.<sup>6,8,9</sup> Dysferlin has a well-studied role in membrane repair, an important process in muscle fibers, which are continually subject to mechanical stress-induced injuries. Mutations in DYSF have been exclusively associated with skeletal muscle diseases. Absence of dysferlin leads to impaired resealing of sarcolemmal wounds.<sup>10</sup> Defects in dysferlin are also known to cause increased inflammatory attack to muscle fibers, which contributes to the exacerbation of the muscle pathology.<sup>11,12</sup>

Currently, there is no treatment for dysferlinopathy. Given that a single gene is causative for the pathology, gene therapy holds great promise. However, the large size of the DYSF coding sequence represents a challenge for gene transfer approaches, since most viral vectors used in gene therapy have a lower cargo capacity. Sleeping Beauty (SB) transposon is a nonviral genetic tool widely used for stable gene transfer in various cell types.<sup>13</sup> This plasmid-based, bi-component system consists of a transposon DNA sequence and a transposase protein that excises the transposon from the donor plasmid and integrates it into the target genome. The transposon can be engineered to carry any gene of interest. Although the efficacy of transposase-mediated transgene insertion decreases with increasing cargo size,<sup>14,15</sup> the hyperactive SB100X transposase is still capable of integrating large, over 10 kb or even BAC-size DNA.<sup>16</sup> Thus, the SB system is well suited to deliver large sequences, such as the DYSF coding sequence. Most importantly, the SB system has already been used in a clinical setup.<sup>17</sup> We constructed an SB transposon-based vector to deliver the full-length human DYSF cDNA into dysferlin-deficient H2K myoblasts (H2K A/J).<sup>18</sup> H2K myoblasts are conditionally immortalized through expression of the tsA58 thermosensitive SV40 large-T-antigen driven by the H-2Kb promoter.<sup>19</sup> They have extensive proliferative capacity in vitro and can engraft robustly into muscle, offering a proper model to test the feasibility of our therapeutic strategy consisting of stably expressing full-length DYSF using the SB system.<sup>18</sup> H2K A/J myoblasts are derived from dysferlin-null mice, harboring a homozygous Dysf<sup>emd</sup> mutation<sup>20</sup> and the H-2K<sup>b</sup>tsA58 allele.<sup>21</sup> Our strategy involved the optimization of a pretransplantation treatment

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by combining irradiation and cardiotoxin (CTX) injection. Our results demonstrate proper dysferlin reconstitution as well as highly efficient engraftment of engineered myoblasts in dysferlin- and immunodeficient B6.Cg-Dysf<sup>emd</sup> Prkdc<sup>−/−</sup>J (Scid/BLA/J) mice.\(^{22}\)

Results

H2K A/J muscle cells properly express full-length human DYSF following SB-mediated gene transfer

To ensure optimal expression of the therapeutic gene, we chose the synthetic c5-12 (Spc5-12) promoter.\(^{23}\) The Spc5-12 promoter was constructed by random assembly of evolutionarily conserved transcription factor binding sites, providing tissue specificity in adult skeletal muscle. Importantly, the Spc5-12 promoter was shown to drive strong transgene expression in vitro in myoblasts and myotubes\(^{23}\) and in vivo in mouse myofibers.\(^{24,25}\) The size of the Spc5-12 promoter is <400 bp. In our hands, a duplicate of Spc5-12 (2xSpc5-12) regulatory sequence proved to be the most efficient in driving transgene expression in H2K A/J myoblasts. We generated a bicistronic vector, in which the full-length human DYSF cDNA was followed by a GFP reporter. GFP was preceded by an Internal Ribosome Entry Site (IRES) sequence to allow simultaneous translation of both cistrons (pT2-2xSpc5-12-hDYSF-IRES-GFP; short: hDYSF-IRES-GFP) (Figure 1a).\(^{15}\) The engineered cells were selected by FACS sorting for the GFP signal after 11 days. At this time, the background GFP signal deriving from nonintegrated plasmid DNA could no longer be detected (Supplementary Figure S1). Due to the rather low transfection rate of the plasmid DNA could no longer be detected (Figure 1b). We counted 500–1,100 dysferlin-positive fibers/cross sectional area in muscles collected 6 weeks posttransplantation (n = 2) (Figure 2a, middle and right columns). We still detected 400–1,000 dysferlin expressing fibers/cross sectional area in muscles collected 6 weeks posttransplantation (n = 2) (Figure 2b, middle and right columns). Per section, the area of the transplant extended up to 1,000 × 800 µm. As in healthy muscle, dysferlin was mainly expressed at the sarcolemma. No dysferlin-positive fibers were detected in the control muscles that were only irradiated or injected with CTX (not shown).

Muscle regeneration using H2K A/J-DYSF<sup>+</sup> cells appears morphologically normal

We next investigated whether transplants derived from H2K A/J-DYSF<sup>+</sup> myoblasts would properly regenerate muscle. Serial cryosections from transplanted TA muscles were immunostained for dysferlin to identify the transplanted area. H&E (Hematoxylin and Eosin), Gomori Trichrome, and NADH-dehydrogenase stains revealed that the grafted area was sharply demarcated towards the host muscle. There was only a mild increase in connective tissue mass between host- and graft-derived muscle. Grafted muscle displayed all typical features of regeneration, including large variation of fiber size, internalized nuclei in a high percentage of fibers, typical features of regeneration, including large variation of fiber size, internalized nuclei in a high percentage of fibers, loose tissue structure, and lack of inflammatory infiltrates (Figure 3).

H2K A/J-DYSF<sup>+</sup> muscle cells gave rise to multiple Pax7-positive cells

Myonuclei of mature fibers are postmitotic and the de novo nuclei are provided exclusively by myoblast–myoblast or myoblast-to-myobifer fusion. Therefore, only engraftment of a subset of donor-derived cells as satellite cells would ensure contribution to muscle regeneration in pathological circumstances and a long-lasting therapeutic effect.\(^{26}\) Satellite cells can be identified by their expression of the transcription factor Pax7\(^{30}\) and their localization beneath the basal lamina of myofibers.\(^{21}\) We found cells positive for the expression of both Pax7 and dysferlin. A few of them located underneath forming basal lamina structures identified by laminin staining, but many of them had not yet adopted a satellite cell position and were interstitial (Figure 4).
Discussion

Here, we describe a proof of principle of an SB-based gene transfer into mouse myoblasts. In the last decades, several approaches have been taken to restore dysferlin function in dysferlinopathies. In allogenic cell transplantation studies, dysferlin expression was reconstituted in mouse myofibers by transplanting labeled wild-type mouse myoblasts into SJL mice. In addition, expression of human dysferlin could be detected following intramuscular transplantation of human control myoblasts into SCID mice. In gene therapy studies, adeno-associated viral (AAV) vectors have been extensively investigated for dysferlin gene transfer. Due to their limited cargo capacity, AAV vectors cannot deliver the full-length DYSF coding sequence. However, in very elegant approaches using the high intermolecular recombination ability of AAVs, it was possible to transfer the 5′ and 3′ moiety of the DYSF cDNA in two separate vectors, achieving robust full-length dysferlin protein reconstitution in vivo. Dual AAV-mediated dysferlin expression has been shown to persist for >1 year in mouse skeletal muscle. However, AAV vectors are non-self-replicating and long-term treatment of patients with skeletal muscle disorders may require repeated vector administrations, with the possible complication of immune responses against the viral capsid.

The use of lentiviral vectors has enabled the successful delivery of full-length dysferlin into patient-derived CD133+ cells in vitro, but the results in vivo have so far been discouraging. In addition, there have been attempts to use RNA-based strategies like exon skipping or trans-splicing to treat dysferlinopathies. Exon 32 of DYSF might be dispensable for protein function, since a patient harboring a mutation in this exon developed a mild disease phenotype. Successful targeting of exon 32 to induce exon skipping has been shown in vitro using antisense oligonucleotides. Furthermore, our laboratory demonstrated successful spliceosome-mediated trans-splicing of the DYSF pre-mRNA both in vitro and in vivo. Despite of the encouraging in vitro results, in vivo dysferlin protein restoration using RNA-based therapies has remained very limited. Our current work using ex vivo delivery of an SB transposon-based therapeutic construct has provided the most efficient results so far, in terms of engraftment and dysferlin protein restoration of any cell.
therapy or cell-based gene therapy studies in dysferlin-deficient mouse models.

We have transplanted engineered H2K A/J-DYSF+ myoblasts into the TA muscles of Scid/BLA/J mice. We determined engraftment efficiency based on the presence of donor-derived myofibers, expressing dysferlin. We could use dysferlin expression directly to monitor engraftment, as mice carrying a homozygous Dysf<sup>md</sup> allele do not express any dysferlin protein. Accordingly, we did not detect any dysferlin-positive fibers in nongrafted muscles of Scid/BLA/J mice that were uninjured, irradiated only, or had just received CTX.

Immortalized C2C12 mouse myoblasts have been shown to rapidly form tumors when transplanted into TA muscles of mdx nu/nu and nondystrophic beige/nu/Xid mice. In those studies, tumor formation was especially evident in muscles that had been irradiated. H2K myoblasts are only conditionally immortalized and are not expected to form tumors in vivo. Still, spontaneous transformation during in vitro culturing could occur and might result in tumor formation following grafting into an immunosuppressed model. Notably, we did not observe any tumors in the grafted muscles or the surrounding tissues at any of the examined time points.
suggesting that the transplanted pool was free of cells spontaneously transformed in culture.

In our hands, a pretransplantation treatment consisting of a combination of irradiation with CTX injection improved engraftment as compared to irradiation alone. Although it is accepted that inactivation of the endogenous satellite cells by radiation or cryoinjury is beneficial for the engraftment of donor cells, the role of additional injury using myotoxins is rather controversial.26,43 The differences in transplantation efficacy might be partially explained by technical reasons. Also, each disease and engraftment model may require a specific pretransplantation treatment that needs to be optimized individually. In this regard, it is important to note that muscle regeneration only occurs when Pax7-positive satellite cells, the proper stem cells of skeletal muscle, are present in the tissue.44,45 In agreement with this assumption, engraftment and contribution to myofibers of nonmuscle stem cell types, like mesoangioblasts or CD133+ cells, have only been demonstrated in nonirradiated muscles of Scid/BLA/J mice, injured with myotoxins only, where the endogenous satellite cells were intact.37,46 Our results clearly support the positive role of a double injury model. Although formation of functional satellite cells by transplanted myoblasts has been reported,19,47 the detection of multiple Pax7-positive, dysferlin-positive cells in the graft’s area was unexpected because engineered H2K A/J myoblasts had been extensively expanded ex vivo prior to transplantation. Whether donor-derived satellite cells with true stem cell characteristics were established remains unresolved.

Our SB-based approach can theoretically be applied to all loss-of-function mutations in DYSF. The effect of SB-mediated DYSF delivery is more difficult to foresee in dysferlinopathies caused by missense mutations in DYSF. In these individuals, the residual dysferlin protein might be accumulated, resulting in aggregates within the muscle cell.48 In these cases, the suitability of this therapeutic approach must first be assessed in vitro, since overexpression of dysferlin might cause a myopathy itself.49 In principle, SB could be used to deliver genes relevant for other muscular dystrophies as well. SB has several advantageous features compared with other integrating vectors, including a potentially safer, close-to-random genomic integration profile versus lentivirus or other transposon-based systems.50,51 Furthermore, since it is a plasmid-based system, it is simple and inexpensive to manufacture, which represents a substantial benefit for clinical use. Maintenance of human primary myoblast cultures, however, is labor intensive and expensive. On the other hand, it remains to be established how human muscle needs to be prepared in order to accept autologous myoblast transplants. If needed, more gentle or alternative treatments to render patients’ muscles receptive to grafts, while avoiding irreversible tissue damage, should be evaluated. Altogether, our results show nonviral integration of full-length human dysferlin into muscle cells as well as successful, efficient transplantation into skeletal muscle. These
are important advances towards successful autologous cell-based gene therapy of dysferlin-deficient muscular dystrophy. Further efforts are needed to enable translation of our findings into clinical use in patients.

Materials and methods

**Plasmid vectors.** The SB100X expression vector was previously described.15 The SB-based reporter vector pT2-CAG-IRES-GFP was kindly provided by Angélica García Pérez, Max Delbrück Center for Molecular Medicine, Berlin. The SB-based vector for full-length dysferlin transfer (hDYSF-IRES-GFP) was constructed as follows: the full-length human dysferlin coding sequence (accession #DQ267935) was extracted from the pDONR221:15803 plasmid (provided by The Jain Foundation) with SpeI + NotI and cloned into an empty SB transposon plasmid previously digested with XbaI + NotI (resulting vector - pT2-hDYSF). The Spc5-12 promoter sequence was extracted from pAAV-Spc5-12-huDysco (Kindly provided by George Dickson, Royal Holloway
University of London) with XbaI. Two copies of the Spc5-12 sequence were inserted in tandem in a 5′ to 3′ orientation into pT2-2xHDSF digested with SpeI (resulting vector - pT2-2xSpc5-12-HDSF). The IRES-GFP-poly(A) sequence was PCR-amplified from pT2-CAG-IRES-GFP with Pfu Ultra II Fusion DNA polymerase (Agilent Technologies, Santa Clara, CA) using the forward 5′-ATCTGGCGCCGCTAGACCATCC GCCCCCT-3′ and reverse 5′-TCTGGCGCCGGCTGTA CACTAGTCGATCCCTCTGTTAAGTACCAC-3′ primers. The resulting PCR fragment was digested with NotI and inserted into pT2-2xSpc5-12-HDSF previously digested with NotI.

Cell culture. Conditionally immortalized H2K and H2K A/J myoblasts were a gift from Terence Partridge, National Children’s Hospital, Bethesda, MD. Cells were cultured on dishes coated with 0.1% gelatin from porcine skin (Sigma-Aldrich, Saint Louis, MO). H2K myoblasts were routinely tested for mycoplasma and experiments were only performed using mycoplasma-free cells. To allow proliferation, H2K myoblasts were cultured at 33 °C and 10% CO2 in Dulbecco’s Modified Eagle Medium high glucose supplemented with GlutaMAX and pyruvate (Gibco, Life Technologies), 20% fetal calf serum (PAA Laboratories, Pasching, Austria), 2% chick embryo extract (US Biological, Salem, MA), and 200 U/ml of γ-interferon (Merck Millipore, Darmstadt, Germany). For differentiation, cells were placed into 37 °C and 5% CO2 for 7 days in proliferation media devoid of γ-interferon and subsequently switched into Dulbecco’s Modified Eagle Medium containing Antibiotic-Antimycotic (Gibco, Life Technologies), 20% fetal calf serum (PA Laboratories, Pasching, Austria), 2% chick embryo extract (US Biological, Salem, MA), and 200 U/ml of γ-interferon. H2K A/J myoblasts was done using a FACSAria flow cytometer (BD Biosciences). Transfection and sorting of H2K A/J myoblasts were transfected with a Neon Transfection System (Life Technologies). An electroporation protocol consisting of 1,050 mV, 30 ms, 2 pulses was selected due to an optimal ratio between transfection efficiency and cell viability. Thoroughly washed cells were aliquoted according to the number of cells needed for each transfection. Each aliquot was resuspended in buffer R (Neon Transfection System, Life Technologies) containing plasmid DNA and electroporated according to the manufacturer’s instructions. Immediately after electroporation, cells were plated in gelatin-coated dishes containing H2K myoblast proliferation media. Routine flow cytometry to monitor expression of GFP was performed using a FACSCalibur flow cytometer (BD Biosciences, Franklin Lakes, NJ) and analysis of the flow cytometry data was performed with the Cellquest Pro (BD Biosciences) and FlowJo Softwares. Sorting of GFP expressing H2K A/J myoblasts was done using a FACSAria flow cytometer (BD Biosciences).

Sodium dodecyl sulfate–polyacrylamide gel electrophoresis and immunoblot. For immunoblotting, myoblasts or myotubes were lysed on ice with radioimmunoprecipitation assay buffer (50 mmol/l Tris–HCl, 150 mmol/l NaCl, 0.1% NP-40, 0.1% sodium dodecyl sulfate, 0.1% sodium deoxycholate, and protease inhibitors). The protein concentration was determined using BCA Protein Assay Kit (Thermo Fisher Scientific, Waltham, MA). For each sample, 10 µg protein diluted in sample buffer (0.25 mol/l Tris–HCl, 50% glycerol, 5% sodium dodecyl sulfate, 0.05% bromophenol blue, and 10% freshly added β-mercaptoethanol) were loaded onto a Novex 8–16% gradient Tris–glycine Mini Protein Gel (Life Technologies). Proteins were separated in denaturing conditions and transferred to a polyvinylidene difluoride membrane using a wet electroblotting system (Bio-Rad, Hercules, CA) in Tris–glycine buffer containing 10% methanol and 0.02% sodium dodecyl sulfate. Blocking was performed with 5% dry milk powder. The membrane was then incubated with the primary antibodies (ab) diluted in blocking buffer (anti-dysferlin, Novocastra NCL-Hamlet, 1:500, overnight at 4 °C) followed by extensive washing. Incubation with the corresponding HRP-conjugated secondary ab was performed at RT for 1 hour. After washing, the membrane was incubated with ECL reagent (GE Healthcare, Milwaukee, WI) for 5 minutes and imaged using a STELLA 3200 system (Raytest, Straubenhardt, Germany). Images were processed using Adobe Photoshop CS5. Any adaptations were applied to the full image with all lanes only.

Dysferlin immunostaining in cultured myoblasts and myotubes. For immunofluorescence staining, H2K myoblasts in ibidi wells (ibidi, Westerburg, Germany) were fixed for 5 minutes in methanol at –20 °C, washed with phosphate-buffered saline (PBS), and blocked with 1% bovine serum albumin/PBS for 1 hour at RT. Dysferlin was detected using the Umbra ab specific to the N-terminal part of dysferlin (Abcam ab124684, 1:200, Cambridge, UK). After washing, a Cy3-conjugated secondary anti-rabbit ab was added (Jackson Immunoresearch 711-165-152, 1:500, West Grove, PA). Nuclei were counterstained with 4′,6-diamidino-2-phenylindole (1:10,000 in PBS). The wells were washed, sealed, and kept at 4 °C until imaging.

Mouse experiments. B6.Cg-Dystmdm Prkdcm/Scid/BLA/J (Scid/BLA/J) homozygous for the Dystmdm and Prkdcm alleles were purchased from The Jackson Laboratory and bred in our specific pathogen-free animal facility. All animal experiments were performed under the license number G0035/14 (LaGeSo, Berlin, Germany). Focal irradiation of mouse hind limbs was performed as described.25 For transplantation, cultured H2K myoblasts were washed and resuspended in Dulbecco’s Modified Eagle Medium containing 2% fetal calf serum. Eight 12–15-week-old female Scid/BLA/J mice were anesthetized with ketamine-xylazine (9 mg/ml ketamine, 1.2 mg/ml xylazine) in sterile PBS at a dose of 160 µl/20 g. The area above the TA was shaved and disinfected and 30 µl of cell suspension were injected with a 26-gauge hypodermic needle in the median portion of the TA.
When CTX was applied, 40 µl of 10 µmol/l CTX/PBS (Latoxan, Valence, France) was injected 1–2 minutes after the cells. At the indicated time points, mice were sacrificed. TA muscles were prepared and cut in two halves following a transversal plane. With the cutting edge atop, each half was separately mounted on cork plates and frozen in liquid nitrogen under cryoprotection. Frozen muscles were stored at −80 °C.

**Immunofluorescence staining of muscle sections.** For dysferlin immunostaining, 6-µm transversal cryosections were air-dried for at least 30 minutes at RT and fixed for 5 minutes in acetone at −20 °C. After blocking, they were incubated overnight at 4 °C with the Romeo ab (1:100) in 1% bovine serum albumin/PBS. Secondary anti-rabbit Cy3-conjugated was applied as above. Nuclei were counterstained with 4',6-diamidino-2-phenylindole (1:5,000). The sections were washed in PBS and ddH₂O and mounted with Aqua Poly/Mount (Polysciences, Inc, Warrington, PA). For Pax7/ laminin immunostaining, 6-µm cryosections were fixed in 4% paraformaldehyde, washed, and blocked with 5% bovine serum albumin, 3% donkey serum. After overnight incubation at 4 °C with a monoclonal mouse-anti-Pax7 ab (DSHB, Iowa City, IA, supernatant diluted 1:1 in sterile glycerol, working dilution 1:10) and rabbit polyclonal anti-laminin ab (Sigma-Aldrich L-9393, 1:200), sections were washed and incubated with secondary abs against mouse IgG (AlexaFluor 555, life technologies, 1:500) and rabbit IgG (AlexaFluor 647, life technologies, 1:500). Counterstain of nuclei, washing and mounting was performed as above. For Pax7/dysferlin immunostaining, 6-µm cryosections were fixed with 4% paraformaldehyde and blocked with 5% bovine serum albumin + 3% donkey serum. The sections were incubated overnight at 4 °C with mouse-anti-Pax7 ab (DSHB, stock as above, working dilution 1:5) and Romeo ab (1:50). Incubation with secondary antibodies against mouse and rabbit IgG as well as nuclei counterstaining and mounting were performed as above. H&E, Gomori Trichrome, and NADH-dehydrogenase stainings were performed according to standard protocols.

**Image acquisition and processing.** All image-containing figures were built in Adobe Illustrator CS5 and image graphics were added in Adobe Illustrator CS5. Images from immunofluorescence stainings on cultured cells and mouse muscle cryosections were acquired with a Leica DM LB2 microscope (Leica Microsystems, Wetzlar, Germany) and processed with Adobe Photoshop CS5. All modifications were always applied to the full image.

**Supplementary material**

**Figure S1.** SB transposition in H2K A/J myoblasts.

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