Abstract. Spinal muscular atrophy (SMA) is a lethal childhood neurodegenerative disorder that is caused by the homozygous deletion of survival motor neuron 1 (SMN1). To date, no effective treatments are available. In the current study, urine cells taken from SMA patients were cultured and the application of patient-derived urine cells was determined in drug intervention. A total of 13 SMA patient-derived urine cell lines and 40 control cell lines were established. SMN was highly expressed in the nucleus and cytoplasm. Patient-derived urine cells expressed low levels of SMN protein compared with controls, they exhibited good tolerance to chemical and electrical damage. SMN expression was upregulated following treatment with histone deacetylase inhibitors and the effect was greater in groups treated with morpholino modified antisense oligo, which targets ISS-N1 in SMN2 intron 7. The results of the current study indicated that SMA patient-derived urine cells may be useful in the initial screening of potential compounds and drugs to treat SMA.

Introduction

Spinal muscular atrophy (SMA) is a lethal neurological genetic disorder that occurs in infancy. There are three types of childhood onset SMA, classified according to age of onset and motor function: SMA I, II and III (1). Patients with SMA I, the most severe type, usually develop muscular weakness before reaching 6 months and succumb to respiratory failure before the age of 2. In mainland China, the prevalence of SMA is ~1.4 in 10,000 and the carrier frequency is estimated to be 1 in 42 (2). To date, no effective treatments for SMA are available.

The SMA-determining gene, survival of motor neuron 1 (SMN1) was initially identified in 1995 (3). SMN2, a highly identical homolog to SMN1 in 5q13 has also been identified, encoding a majority of the truncated SMN protein (Δ7-SMN) that results from the abnormal splicing of exon 7 during transcription (3). The SMN2 gene is unique to humans and modifies the disease severity of SMA, thus representing a promising therapeutic target. Different types of drugs stimulating the inclusion of exon 7 in SMN2 have been identified, including histone deacetylase inhibitors (HDACi), hydroxyurea, ceftriaxone, antisense oligo and novel synthetic compounds (4-6). The efficacy of valproic acid has been assessed in clinical trials, however the results are controversial, as both positive and negative results were observed in SMA patients from different clinical trials (7-11).

SMA is a genetic disease, thus the acquisition of SMA patient-derived cell samples may function as a tool for molecular research and drug intervention. However, although patient-derived fibroblasts are currently used widely in research to assess the mechanism of a number of neurological disorders, muscle or skin biopsy procedures are invasive and usually unacceptable for young patients with SMA clinically. Previously, urine cell lines have been successfully established from urine sediments (12). In the present study, urine sediments from different patients with SMA were cultured and patient-derived urine cell lines were established in vitro. Using these stable urine cell lines, HDACi and antisense oligo intervention was performed, aiming to investigate the application of urine cells in screening the efficacy of potential drugs to treat SMA.

Materials and methods

Primary urinary cell culture. Urine sediment culture was performed following a previously described protocol (12). All recruited patients fulfilled the clinical and genetic diagnostic criteria for SMA with homozygous deletion of SMN1 gene (13). A total of 13 patients with SMA (12 males and 1 female; age range, 1.5-39 years) were recruited in the
current study between June 2011 and September 2013 from the First Affiliated Hospital of Fujian Medical University (Fuzhou, China). A total of 40 control urine cell lines were cultured, using the same culture method, from control subjects (36 males and 4 females, aged 5-62 years) without SMA disease at the same period (June 2011 to September 2013) from the First Affiliated Hospital of Fujian Medical University (Fuzhou, China). The present study was approved by the Ethics Committee of First Affiliated Hospital of Fujian Medical University and written informed consent was obtained from all participants or their parents.

Valproic acid (VPA) and Suberoylanilide hydroxamic acid (SAHA) intervention. A total of 13 SMA urine cell lines were produced from different patients. The majority of the urine cell lines consisted of fusiform cells with similar cell growth rates. The current study used 50 mM VPA and 20 µM SAHA to investigate the toxicity of VPA and SAHA in urine cells, cell toxicity analysis to assess the rate of cell death. Experiments were repeated at least three times.

After 6 h, the medium was switched to fresh epithelial cell medium (ScienCell Laboratories, Inc.) at 37˚C for 6 h. Cells were seeded onto 12-well plates with 3x10^4 cells/well. Following electroporation, urine cells were treated with 50.0 msec; Capacity (Capa), 1416.3 uF; Pp off, 10.0 msec and electroporation were: Poration pulse (Pp) V , 150 V; Driving voltage (Vd) 200 V; and capacitor (Cp) 0.18 µF; Pulse off time (Pp off), 50.0 msec. Following electroporation, urine cells were split the cells on ice for 10 sec. Lysis buffer was collected (20 µl per lane) were separated using 12% SDS-PAGE and incubated for 24, 48 and 72 h, morphological changes in the cells were observed and SMN expression was quantified. All experiments were repeated at least three times. The concentration of VPA and SAHA was adopted according to previous studies (14,15).

Morpholino modified antisense oligo (ASO) intervention. A previous study observed that morpholino-ASO was able to significantly increase the expression of SMN protein (16). Therefore, morpholino-ASO was purchased from Gene Tools, LLC, Philomath, OR, USA). The morpholino-ASO sequence was attcacttttcataatgtctgg, targeting intronic splicing silencer N1 (ISS-N1) in SMN2 intron 7. After 6 h, the morpholino-ASO was added and seeded onto 96-well plates with 2-3x10^4 cells/well. When cell density reached ~80%, VPA and SAHA were added and incubated for 24 h, 48 h and 72 h. At each different time point, cells were fixed with 2.5% formaldehyde (at room temperature for 30 min) and permeabilized with 0.1% Triton-X-100 (at room temperature for 5 min) in phosphate-buffered saline (PBS). Following blocking with 5% non-fat milk (at room temperature for 1 h), cells were incubated with anti-SMN1 rabbit polyclonal antibody (1:500; GTX101047; Genetex, Inc., Irvine, CA, USA) at 4˚C overnight and were subsequently probed with peroxidase-conjugated goat anti-rabbit immunoglobulin G (IgG; ZF-0516, ZSGB-BIO Technology, Co., Ltd., Beijing, China) at 37˚C for 2 h. Tetramethyl benzidine developing solution (Promega Corporation, Madison, WI, USA) was added and 15 min later, the reaction was attenuated using 2 mol/l H_2SO_4. Optical density values were measured at 490 nm using an ELISA reader (Bio-Rad, Laboratories, Inc., Hercules, CA, USA) and were further analysed using Excel software (Office, Version 10.1.0.6393; Microsoft Corporation, Redmond, WA, USA).

SMN-specific protein cell immunoassay. The cell immunoassay was performed as previously described by Sumner et al (17), with minor modifications. Briefly, urine cells were digested and seeded onto 96-well plates with 2-3x10^4 cells/well. When cell density reached ~80%, VPA and SAHA were added and incubated for 24 h, 48 h and 72 h. At each different time point, cells were fixed with 2.5% formaldehyde (at room temperature for 30 min) and permeabilized with 0.1% Triton-X-100 (at room temperature for 5 min) in phosphate-buffered saline (PBS). Following blocking with 5% non-fat milk (at room temperature for 1 h), cells were incubated with anti-SMN1 rabbit polyclonal antibody (1:500; GTX101047; Genetex, Inc., Irvine, CA, USA) at 4˚C overnight and were subsequently probed with peroxidase-conjugated goat anti-rabbit immunoglobulin G (IgG; ZF-0516, ZSGB-BIO Technology, Co., Ltd., Beijing, China) at 37˚C for 2 h. Tetramethyl benzidine developing solution (Promega Corporation, Madison, WI, USA) was added and 15 min later, the reaction was attenuated using 2 mol/l H_2SO_4. Optical density values were measured at 490 nm using an ELISA reader (Bio-Rad, Laboratories, Inc., Hercules, CA, USA) and were further analysed using Excel software (Office, Version 10.1.0.6393; Microsoft Corporation, Redmond, WA, USA).

Western blotting. Urine cells were seeded onto 6-well plates at a density of 3-5x10^4 cells/well with epithelial cell medium at 37˚C for 24 h. When cell confluence reached ~80%, total proteins were extracted by cell lysis buffer for western blot (catalogue no. P0013; Beyotime Institute of Biotechnology). Briefly, 100 µl cell lysis buffer was added to each well to split the cells on ice for 10 sec. Lysis buffer was collected and centrifuged at 12,000 x g for 5 min at 4˚C to harvest the supernatant. The supernatant was subsequently denatured at 100˚C in water for 10 min. The protein concentration was determined using the BCA protein assay kit (catalogue no. P00125; Beyotime Institute of Biotechnology) following manufacturers protocol, using a microplate reader (SpectraMax M5; Olympus Corporation, Tokyo, Japan) according to the manufacturer’s protocol. Proteins (20 µl per lane) were separated using 12% SDS-PAGE and transferred to nitrocellulose membranes (GE Healthcare, Chicago, IL, USA). The membrane was blocked at room temperature for 1 h using 5% skimmed milk in Tris-HCl buffer. It was then probed with rabbit polyclonal SMN1 antibody (1:500; catalogue no. GTX101047; Genetex, Inc.) and mouse monoclonal Actin antibody (1:20,000; catalogue no. AA128; Beyotime Institute of Biotechnology overnight at 4˚C. Peroxidase-conjugated goat anti-rabbit (catalogue no. ZB-2301; ZSGB-Bio, Technology, Co., Ltd.) and mouse

Cell toxicity analysis to assess the rate of cell death. To investigate the toxicity of VPA and SAHA in urine cells, the current study used 50 mM VPA and 20 µM SAHA to treat urine cells and observed the cell morphological changes and cell mortality rates using a Trypan blue staining cell viability assay kit (catalogue no. C0011; Beyotime Institute of Biotechnology, Shanghai, China) following the manufacturers protocol. The number of trypan blue positive cells was calculated using a hemocytometer (Shanghai Precision Instruments Co., Ltd, Shanghai, China) under an inverted microscope (IX71; Olympus Corporation, Tokyo, Japan). The ratio of trypan blue positive cells to total cells when seeded was then calculated.
IgG (ZB-2305, ZSGB-Bio, Technology, Co., Ltd.) were used as secondary antibodies at a dilution of 1:2,000 at 37˚C for 2 h. Images were acquired using darkroom development techniques for chemiluminescence (BeyoECL Plus A and B solution; Beyotime Institute of Biotechnology). The ratio of SMN to Actin was calculated using QuantityOne software, version 4.6.2 (Bio-Rad Laboratories, Inc.).

**Immunofluorescence staining.** Urine cells (SMA -01 and control -01) were seeded onto cover slips in a 6-well dish (1x10⁴ cells/slip). The adherent cells were then fixed with 4% paraformaldehyde (at room temperature for 30 min) and were blocked with 10% goat serum in PBS (at room temperature for 1 h). Cells were then probed using a SMN1 mouse monoclonal antibody (1:50; catalogue no. sc-32313; Santa Cruz Biotechnology, Inc., Dallas, TX USA) at 4˚C overnight. Alexa Fluor 594 conjugated affinipure goat anti-mouse IgG (H+L) (1:200; catalogue no. ZF-0513; ZSGB-Bio, Technology, Co., Ltd.) was used as the secondary antibody incubated at room temperature for 2 h. Cell nuclei were counterstained with 4', 6-diamidino-2-phenylindole. The cover slips were mounted and visualized using a confocal microscope (Leica TCS SP5; Leica Microsystems GmBH, Wetzlar, Germany).

The current study established 13 urine cell lines from 13 different patients with SMA (Table I) and 40 urine cell lines from controls. SMN protein was highly expressed in the nucleus and cytoplasm both in SMA and control urine cells (Fig. 1A and B). Total levels of SMN protein were determined by western blotting (Fig. 1C), which demonstrated that urine cells derived from patients with SMA expressed significantly lower levels of SMN compared with controls (t=7.522, P=0.002; Fig. 1D).

**Statistical analysis.** One-way ANOVA followed by least significant difference post hoc analysis was performed and assessed using SPSS version 16.0 (SPSS, Inc., Chicago, IL, USA) to analyse the SMN protein variation between drug-treated and control groups. P<0.05 was determined to represent statistically significant differences.

**Results**

**SMN protein expression and localization in urine cells.** The current study established 13 urine cell lines from 13 different patients with SMA (Table I) and 40 urine cell lines from controls. SMN protein was highly expressed in the nucleus and cytoplasm both in SMA and control urine cells (Fig. 1A and B). Total levels of SMN protein were determined by western blotting (Fig. 1C), which demonstrated that urine cells derived from patients with SMA expressed significantly lower levels of SMN compared with controls (t=7.522, P=0.002; Fig. 1D).

**HDACi intervention on urine cells.** To determine the application of urine cells in drug interventions, small molecular compounds known as HDACi, including VPA and SAHA were used. Cell death were observed in groups treated with 50 mM VPA (54±7%) and 20 µM SAHA (49±7%), indicating that high doses of HDACi induce toxicity in vitro. At concentrations of 0, 5, 10, 15 and 20 mM, VPA did not increase levels of SMN significantly after 24, 48 and 72 h treatment (Fig. 2A-C). By contrast, levels of SMN significantly increased following 24 h treatment with 0.5, 1, 5 and 10 µM SAHA (all P<0.05);
however, the drug-induced effects of SAHA were reversed following the longer treatment times of 48 and 72 h (Fig. 2D). Expression of SMN increased significantly in the group treated with 1.0 µM SAHA (P<0.05; Fig. 2E and F). Similar VPA and SAHA intervention results were also observed in SMA-02, SMA-03 and SMA-13 urine cell lines (data not shown).

Morpholino-ASO intervention in urine cells. Morpholino-ASO interventions on urine cells (SMA-01 and SMA-13 lines) were performed using an electroporator. The majority of urine cells were sustained in stable cell morphology following electrical trauma, indicating that the urine cells were able to tolerate physical damage (Fig. 3A and B). Compared with the control group (0 pmol), levels of SMN expression significantly increased by 51.0, 51.4 and 94.6% in the groups treated with 10, 20 and 40 pmol morpholino-ASO, respectively in the SMA-01 cell line (all P<0.05; Fig. 3C and D). Similar ASO intervention results were observed in the SMA-13 cell line (data not shown).

Discussion

In the present study, 13 SMA patient-derived non-invasive urine cell lines were successfully cultured. The urine cells grew robustly in vitro, carried SMN gene mutations (12) and expressed low levels of SMN. Furthermore, they exhibited tolerance to small molecular compounds and electrical trauma, demonstrating that they may be used effectively in drug interventions.

There have been large advances in the field of SMA therapy, including the use of small-molecule drugs, RNA-based therapy, SMN gene correction and stem cell-based cell replacement (1,18). Previous studies have demonstrated that as a first-generation HDACi, VPA is able to increase SMN2 expression in SMA patient-derived fibroblasts, reduce the degeneration of motor neurons and increase motor function in mice with SMA (14,19,20). In a clinical trial, Weihl et al (7) reported that 7 patients with type III/IV SMA who were treated with VPA experienced an increase in muscle strength and motor function. However, in several subsequent large clinical trials, VPA or VPA + L-carnitine failed to improve the muscle strength or motor abilities of patients with SMA (8-11). SAHA is an orally administered HDACi used to treat cutaneous T cell lymphomas that was approved by FDA in 2006 (21). As a second-generation HDACi, SAHA may suppress type I and type II HDACi, while VPA only suppresses type I HDACi selectively (22). SAHA may activate the SMN2 gene more effectively than VPA and ameliorate the SMA phenotype in mouse models of SMA (15,23,24). The results of the current study further supported the hypothesis that SAHA appears to be more efficient than VPA in increasing SMN protein expression in urine cells.

ASO, a type of RNA-based therapy, is able to significantly increase expression of SMN. ASO can effectively block...
targeted exon splice enhancers (ESEs) or intronic splicing silencers (ISSs) within SMN2, including element 1, exonic splicing silencers A, exonic splicing silencers B, ISS-N1 and ISS+100 (25,26). The negative regulator ISS-N1 is the most effective at increasing the expression of SMN (16,26). Previous studies have demonstrated that peripheral and intracerebroventricular administrations of ASO may render the inclusion of exon 7, upregulate the SMN protein expression, rescue the motor neuron function, and extend the lifespan in SMA mouse model (27,28). ASO interventions are also effective at the age of onset, even following the onset of neurological symptoms in SMA mice (29). Few studies have been performed on cell samples derived from patients with SMA. Thus the current study performed ASO intervention in urine cells taken from patients with SMA, and demonstrated that ASO may significantly upregulate SMN expression.

The primary limitation of the current study is that urinary epithelial cells are a non-neural cell type. However, as a type of patient-derived somatic sample, urine cells carry SMN gene mutations and express a relatively low level of SMN protein. Compared with obtaining patient-derived neurons, the urine cell culture process is simple and non-invasive; thus urine cells may be used as a platform for drug intervention and provide initial evidence indicating whether a candidate drug to treat SMA is able to upregulate the expression of SMN in cells derived from patients with SMA.

In conclusion, the results of the current study indicate that urine cells derived from patients with SMA exhibit tolerance to chemical and physical damage, which may be useful in the initial screening of potential compounds and drugs to treat SMA.

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