**Tm7sf2 may participate in the healing of burn wounds**

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**Abstract.** The transmembrane 7 superfamily member 2 (Tm7sf2) gene has been reported to be involved in cholesterol biosynthesis by encoding the protein 3β-hydroxysterol A14-reductase (8). Tm7sf2 participates in the cellular response to stressful conditions (9) and its loss alters the expression of proteins that are involved in the differentiation of epidermal cells by reducing cholesterol sulfate levels (10). The loss of the entire epidermis following burn injury may induce severe hematopoietic dysfunction, which will subsequently increase the risk of mortality and morbidity (11). A previous study indicated that cholesterol levels of the erythrocyte membrane in patients with large area burns were significantly increased. Therefore, it is possible that Tm7sf2 may be involved in the wound healing process (12).

The transmembrane 7 superfamily member 2 (Tm7sf2) gene has been reported to be involved in cholesterol biosynthesis by encoding the protein 3β-hydroxysterol A14-reductase (8). Tm7sf2 may participate in the healing of burn wounds (12). The transmembrane 7 superfamily member 2 (Tm7sf2) gene has been reported to be involved in cholesterol biosynthesis by encoding the protein 3β-hydroxysterol A14-reductase (8). Tm7sf2 may participate in the healing of burn wounds (12).

**Introduction**

Burn injury is one of the most common and severe trauma morphologies and is typically established from thermal injury of the skin (1). The clinical features of burn injuries range from difficult initial assessment to the tendency of pathological scars (2). According to a previous 33-year population-based study, adjusted for all-cause mortality, the mortality rate of burn injury was 1.6 (95% confidence interval, 1.3-2.0), and children with burn injuries had a 1.6-fold higher mortality rate than those without any injuries (3). The resuscitation of burn trauma patients remains a challenge, as the healing mechanism remains to be fully elucidated. When treating burn wounds it is crucial to restore and close the exposed wound surface promptly, as this will aid in limiting the loss of water, electrolytes and nutrients, and preventing the invasion of pathogenic microorganisms (4-6). Aspects of the wound healing mechanism remain unclear; however, it has been established that multiple cell types, the extracellular matrix and soluble mediators are all involved in this process (7).

**Materials and methods**

**Burn wound model.** Sprague-Dawley rats (age, 5 weeks old; male and female; n=10) weighing between 250 and 300 g and maintained under controlled temperature (20±1°C), humidity (55±10%) and illumination (12 h dark:light), were subjected to limb electrical burns as described previously (14). Subsequent
to undergoing anesthesia with 3% pentobarbital (10 mg/kg; Sigma-Aldrich, St. Louis, MO, USA), the rats were maintained in a stationary position on the experimental table in order to shave the hip and right posterior limb. Next, a null electrode was placed on the right limb (0.5 cm proximal to ankle), with a live electrode placed on the ipsilateral gluteus. The electricity (output voltage: 1,000 V) was administered for 0.1 sec to conduct the burns using the YLS-5Q Skin Burning Device model (Biowill Co, Ltd. Shanghai, China). Blood samples were obtained from the tail vein (using 1 ml syringe) at 6 and 24 and 36 h, subsequent to the administration of the electric burns, and then centrifuged at 300 x g, -4˚C for 15 min, to obtain the blood serum.

**Cell culture.** HaCaT cells (Shanghai Bogoo Biotechnology Co., Ltd., Shanghai, China) were cultured in Dulbecco’s modified Eagle’s medium (DMEM; Gibco; Thermo Fisher Scientific, Inc., Waltham, MA, USA) medium supplemented with 10% (v/v) fetal bovine serum, 100 U/ml streptomycin and 100 U/ml penicillin at 37˚C in a 5% (v/v) CO₂ humidified atmosphere. DMEM (Gibco; Thermo Fisher Scientific, Inc.) containing 0.5 mg/ml collagenase type I was applied at 37˚C for 3 h to separate fibroblasts from rat blood serum. These fibroblasts were pelleted by centrifugation at 300 x g, for 7 min, at room temperature, and then cultured in the aforementioned conditions. Cells passaged 2-4 times were used for further experiments.

**Tm7sf2 transfection.** The full-length wild-type Tm7sf2 coding sequence (alternate names: D14SR, SR-1; RefSeq accession, NP_003264) was sub-cloned into pcDNA3.1’ to produce the Tm7sf2 expression vector (pcDNA3.1 Tm7sf2), which was confirmed by sequencing at Shanghai Majorbio Bio-pharm Technology Co., Ltd. (Shanghai, China). Tm7sf2-specific siRNA and control siRNA were synthesized by Shanghai GenePharma Co., Ltd. (Shanghai, China). Lipofectamine 2000 reagent (Invitrogen; Thermo Fisher Scientific, Inc.) was used for the cell transfection, according to manufacturer's protocol, and empty construct pcDNA3.1 was also transfected as a control. Stable Tm7sf2 transfectants were selected using G418 (Gibco; Thermo Fisher Scientific, Inc.).

**Cell interference experiments.** HaCaT cells were divided into four groups and exposed to the following treatments for 20 h: i) Control, 20% normal serum; ii) burn, 20% electrical burned serum; iii) 20% normal serum + siTm7sf2; iv) 20% electrical burned serum + siTm7sf2. The full-length wild-type Tm7sf2 expression vector (pcDNA3.1 Tm7sf2), which was confirmed by sequencing, was sub-cloned into pcDNA3.1’ to produce the Tm7sf2 expression vector (pcDNA3.1 Tm7sf2), which was confirmed by sequencing at Shanghai Majorbio Bio-pharm Technology Co., Ltd. (Shanghai, China). Tm7sf2-specific siRNA and control siRNA were synthesized by Shanghai GenePharma Co., Ltd. (Shanghai, China). Lipofectamine 2000 reagent (Invitrogen; Thermo Fisher Scientific, Inc.) was used for the cell transfection, according to manufacturer's protocol, and empty construct pcDNA3.1 was also transfected as a control. Stable Tm7sf2 transfectants were selected using G418 (Gibco; Thermo Fisher Scientific, Inc.).

**Cell proliferation and invasiveness.** A 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide (MTT; Sigma-Aldrich) colorimetric assay and Matrigel (BD Biosciences, Franklin Lakes, NJ, USA) invasion chamber assay were used for the determination of cell proliferative and invasive capacities, respectively, as described previously (15,16). Each experiment was performed three times.

**Monocyte-endothelial cell adhesion assay.** Calcein-acetomethyl (25 mg; Molecular Probes; Thermo Fisher Scientific, Inc., in 5 ml dimethyl sulfoxide/ml media; Sigma-Aldrich) was used for labeling the treated HaCaT cells. Subsequent to centrifugation at 300 x g, for 7 min, at room temperature, the cells were washed three times with phosphate-buffered saline (PBS; Sigma-Aldrich) to allow adherence to a compact monolayer of the EA.hy926 human endothelial cell line (Cell bank of Type Culture Collection of the Chinese Academy of Sciences, Shanghai, China) at 37˚C for 1 h. Subsequently, the monocytes in the medium were aspirated. A fluorescence microscope (Olympus CX31; Olympus Corporation, Tokyo, Japan) was used for the visualization of cell adherence at excitation and emission wavelengths of 488 and 555 nm, respectively. Six fields were selected per well to calculate the number of adherent monocytes using Image J software (version 2; National Institutes of Health, Bethesda, MD, USA).

**Reverse transcription-quantitative polymerase chain reaction (RT-qPCR).** RT-qPCR was performed as previously described (17). Total mRNA was isolated from cells, and complementary DNA (cDNA) was produced using reverse transcriptase (iScript cDNA Synthesis kit; Bio-Rad Laboratories, Inc., Hercules, CA USA). Tm7sf2 expression levels were measured via SYBR-Green based qPCR (SYBR Green Master mix; Thermo Fisher Scientific, Inc.). Reactions were performed with the initial denaturation at 95˚C for 5 min and then 40-50 cycles at 95˚C for 30 sec, 58-62˚C for 30 sec and 72˚C for 30 sec, and a final extension at 72˚C for 10 sec.

**Western blot analysis.** A 10-12% sodium dodecyl sulfate-polyacrylamide gel (Thermo Fisher Scientific, Inc.) was used for protein separation at 100 V for 90 min. Next, the proteins were blotted onto polyvinylidene difluoride membranes, which were blocked with PBST (0.1% Triton X-100; Sigma-Aldrich in PBS), and probed with primary antibodies including anti-Tm7sf2 (cat no. sc-162325) Beclin 1 (cat no. sc-11427) and LC3-II (cat no. sc-28266) all diluted 1:200 in bovine serum albumin and horseradish peroxidase-conjugated secondary antibodies (sc-391122; diluted 1:5,000) and GAPDH (sc-59540l; diluted 1:1,000). GAPDH was used as a loading control. All antibodies were purchased from Santa Cruz Biotechnology, Inc., (Dallas, TX, USA) unless otherwise stated. Enhanced chemiluminescence (GE Healthcare Life Sciences; Uppsala, Sweden) was used for detecting the bands. The results were quantified using Image J software (version 2; National Institutes of Health, Bethesda, MD, USA).

### Table I. Tm7sf microRNA and protein expression levels.

<table>
<thead>
<tr>
<th>Time (h)/group</th>
<th>mRNA level</th>
<th>Protein level</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>1.00±0.00</td>
<td>-</td>
</tr>
<tr>
<td>6</td>
<td>1.18±0.07</td>
<td>-</td>
</tr>
<tr>
<td>12</td>
<td>1.32±0.05</td>
<td>-</td>
</tr>
<tr>
<td>24</td>
<td>1.49±0.09</td>
<td>-</td>
</tr>
<tr>
<td>36</td>
<td>1.25±0.05</td>
<td>-</td>
</tr>
<tr>
<td>Control</td>
<td>-</td>
<td>1.00±0.00</td>
</tr>
<tr>
<td>Tm7sf2</td>
<td>-</td>
<td>1.43±0.11</td>
</tr>
<tr>
<td>siTm7sf2</td>
<td>-</td>
<td>0.51±0.09</td>
</tr>
<tr>
<td>Tm7sf2+siTm7sf2</td>
<td>-</td>
<td>1.03±0.12</td>
</tr>
</tbody>
</table>

mRNA and protein expression levels are presented as the fold-changes from the 0 h and control groups, respectively. Tm7sf2, transmembrane 7 superfamily member 2; si, small interfering.
utilized to develop the immunoreactive protein bands, which were analyzed by a densitometer (Image Pro Plus, version 1.63; Media Cybernetics, Rockville, MD, USA).

Statistical analysis. Each sample was processed and measured three times. The data are presented as the mean ± standard deviation. The statistical analysis was performed using SPSS software (version 17.0; SPSS, Inc., Chicago, IL, USA) statistical software, one-way analysis of variance, followed by Fisher’s Protected Least Significant Difference post-hoc.
analysis used to determine statistical differences between groups. P<0.05 was considered to indicate a statistically significant difference.

Results

Tm7sf2 level detection. Compared with the control group, Tm7sf2 mRNA levels in rats with electric burns were significantly higher with the increasing time following the administration of the burn (P<0.05). The levels were then reduced after 24 h (Fig. 1A). In the 20% electrical burned serum group, the protein levels of Tm7sf2 were significantly higher than the control group (P<0.05). In the siTm7sf2-transfected cells, Tm7sf2 expression was significantly lower compared with the control group (P<0.05), however was restored to normal with the addition of Tm7sf2 (Fig. 1B; Table I).

Cell proliferation. The MTT assay demonstrated that the burn serum significantly reduced cell proliferation when compared with the normal serum (P<0.05). Furthermore, the proliferation of cells treated with Tm7sf2 siRNA was significantly lower in the burn serum group compared with the control cells (P<0.05; Fig. 2; Table II).

Monocyte-endothelial cell adhesion. The number of monocytes represent the adhesion strength between the EA.hy926 and HaCaT cells (18). The 24 h treatment with burn serum induced the highest expression levels of Tm7sf2, therefore cell adhesion of the HaCaT cells was detected subsequent to 24 h of treatment. The burn serum treatment significantly enhanced the capacity of monocyte-endothelial cell adhesion, even if the cells had been transfected with siTm7sf2, compared with normal serum treatment (P<0.05; Fig. 3; Table III).

Autophagy-associated protein levels. To investigate the influence of the burn serum and siTm7sf2 on autophagic abilities of HaCaT cells, the autophagy-associated Beclin1 and LC3-II protein levels in HaCaT cells were detected using western blot analysis. As indicated in Fig. 4, burn serum (treated for 24 h) increased the expression levels of Beclin1 and LC3-II when compared with control groups (P<0.05). The siTm7sf2 group also presented increased expression levels, while the effect was more evident in the burn serum-treated cells (P<0.01; Table IV).

Discussion

In the current study, it was identified that burn serum treatment reduced cell proliferation, increased monocyte-endothelial cell adhesion, and induced higher expression levels of Beclin1 and LC3-II in HaCaT cells. Additionally, the administration of siTm7sf2 had a similar influence on these properties, suggesting that Tm7sf2 may be important during burn wound repair.

Thermal injury induces the release of inflammatory factors (19). Monocyte-endothelial cellular adhesion, as a key feature in inflammation, is important in monocyte extravasation (14). The observations of the present study indicate that silencing of Tm7sf2 and treatment with burn serum increase cellular adhesion, with the siTm7sf2 + burn serum treatment resulting in the highest number of monocytes. Therefore,
Tm7sf2 may be involved in inflammation during the wound healing process.

Autophagy is a catabolic process necessary for the maintenance of the cellular organelle balance, and is a highly conserved pathway for the delivery of intracellular macromolecule waste to lysosomes (20). Severe burns lead to endoplasmic reticulum stress that triggers autophagy signaling cascades. However, it has been identified that there is reduction in autophagy in burn wounds during the initial period of progression of burn injuries (21). The enhanced level of autophagy deep in the dermis has been suggested as a potential prosurvival factor against injury-associated inflammation (22). The Beclin1 protein is associated with the activation of autophagy and in addition, as a reliable biomarker of autophagy activity, it forms part of the phosphatidylinositol 3-kinase complex (23). LC3 is a critical protein for autophagy, and its cleavage leads to the production of LC3-I, which is converted to LC3-II subsequent to lipidation via a ubiquitin-like system that allows LC3 to be involved in autophagic vesicles (24). The LC3-III/LC3-I ratio is considered to be an indicator of autophagy (25). Furthermore, the increased expression levels of Beclin1 and LC3-II, two indicators of autophagy, observed in the current study suggested that the cells were undergoing the wound healing processes. Tm7sf2 also appears to be crucial to these processes.

In conclusion, burn serum extracted from an electrical burn rat model increased the expression of Tm7sf2. In addition, Tm7sf2 may participate in the wound healing process by interacting with LC3-II and Beclin1 and thus, therapeutically targeting Tm7sf2 may be advantageous to the wound healing process following burn injuries. However, only rat models were used in the present study regarding the role of Tm7sf2, and further clinical or human tissue based studies are required to determine the benefit of novel therapies targeting Tm7sf2 for burn wound healing.

References