Effect of “HuChou San” on ABCC11 mRNA levels and MRP8 protein expression in skin tissue from bilateral axilla of axillary osmidrosis patients

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INTRODUCTION
Axillary osmidrosis (AO) is a clinical condition that manifests in an offensive and unpleasant body odour originating from the axillary area. This odour, which is produced by secretions from the axillary apocrine glands, can cause significant distress and impair the psychosocial well-being of the affected individuals. Many methods are used for treating AO. These strategies may be either in the form of drugs or in the form of surgical intervention. The use of...
conservative strategies such as topical application of astringents, systemic agents, or botulinum toxin, results only in transient therapeutic outcomes [1]. Since surgery may provide a permanent solution for the problem, surgical strategies are preferred by patients with more complicated cases of AO [1-4].

Several surgical techniques have been developed for removal of the axillary apocrine glands. These procedures include removal of subcutaneous tissue, total excision of skin and subcutaneous tissue, ultrasonic liposuction, electrodesiccation, as well as sympathectomy. However, these surgical techniques have yielded variable results [5-7]. Although some of these techniques have been shown to be efficacious [1, 8], concerns still remain regarding their invasive tendency, potential complications, prolonged recovery time, and scar tissue potential. These concerns may influence the satisfaction of the affected individuals. Thus, newer surgical techniques have been evolved to overcome these undesirable side-effects. Herbs have also been used for AO therapy. Saponin, Halite violaceous, Rhizoma typhonii, and Impatiens balsam produce good effects on AO. “HuChouSan” has very strong therapeutic effect on AO, based on the results of a previous study. This prompted the present study aimed at further investigations on the efficacy of “HuChouSan”.

The objective of this study was to investigate the effect of daily treatment with “HuChou San” powder on AO. Abnormal expressions of MRP8 and ABCC11 have been linked to AO [9]. A strong positive preference for low-odorant partners with a dysfunctional ABC11 gene seems a plausible explanation for this striking frequency of loss-of-function allele. Thus, the expression levels of ABCC11 mRNA and MRP8 protein were measured in skin tissues from the bilateral axilla of AO patients before and after “HuChou San” treatment.

METHODS

Ethical considerations

This study received approval from the ethical committee of HuaShan Hospital (approval no. H201904) in accordance with the Declaration of Helsinki, promulgated in 1964 and amended in 1996 [10]. All subjects provided written informed consent.

Materials

“HuChou San” powder consisted of saponin (25 %), Halite violaceous (25 %), Rhizoma typhonii (25 %), and Impatiens balsam (25 %). It was prepared by grinding together equal amounts of saponin, Halite violaceous, Rhizoma typhonii, and Impatiens balsam, into a fine powder.

Patients, drugs and sampling

A total of 40 AO subjects (25 men and 15 women) undergoing concurrent treatment at Shanghai Ninth People’s Hospital from July 2015 to July 2016 were selected as subjects for this study. The ages of the subjects ranged from 18 to 35 years (mean age = 23.7 years). The subjects were treated with “HuChou San” powder according to the following procedure: patient axillary regions were initially treated for 10 min with a solution of copper sulphate and zinc sulphate. Then, “HuChou San” powder was applied gently to the affected region once daily for 3 months. Fresh skin tissue samples surgically collected from the bilateral axilla were preserved at -70°C prior to analysis. The tissue specimens were collected from each participant at four time points. The first sample was taken prior to the initial “HuChou San” treatment (0 month sample), while the remaining samples were taken after the 1st, 2nd and 3rd months of daily “HuChou San” treatment.

Ten healthy individuals (without AO) were also selected as untreated, non-AO controls. Tissue samples were collected from the non-AO control patients at a single time point.

Lysate preparation and Western blotting

Lysates of frozen bilateral axillary skin tissue samples were prepared for Western blotting. The tissue samples were rinsed twice in ice-cold PBS, followed by lysing with RIPA buffer (Beyotime, Shanghai, China) containing 1% PMSF (Lullaby Pharmaceutical Chemical Co. Ltd, Wuhan, China). The lysates were ultrasonicated on ice, and subjected to brief centrifugation at 14,000 × g in the cold. The protein contents of the clear lysates were quantified using standard bicinchoninic acid (BCA) method, before denaturation.

Protein samples (10 – 50 μg total protein) were separated on SDS-polyacrylamide gel electrophoresis using 10 or 12 % tris-glycine gels, and transferred to polyvinylidene difluoride (PVDF) membranes (KaiJie Membrane Seperation Technology Co. Ltd, HangZhou, China). The membranes were blocked with 5 % non-fat milk, prior to incubation overnight with primary antibodies targeting MRP8 (ABCC11) (1:200; Santa Cruz Biotechnology, Santa Cruz, CA, USA) and β-actin (1:10,000; Bioss
Antibodies, Beijing, China) at 4°C. This was followed by incubation with goat anti-rabbit IgG (secondary antibody) conjugated to HRP (AmyJet scientific Inc, WuHan, China). Thereafter, the membranes were probed using ECL reagent (Solarbio Science & Technology Co. Ltd., BeJing, China), followed by exposure to X-ray film (Fujifilm, Tokyo, Japan). The signal intensities of the protein bands were quantified using Quantity One software (Bio-Rad, Hercules, CA, USA). The MRP8 protein expression was normalized to that of β-actin which served as internal standard.

Quantitative polymerase chain reaction (qPCR)

Isolation of total RNA from bilateral axillary skin samples was done with RNX-Plus Solution (SinaClon, Iran). Using 1 μg extracted RNA, cDNA was synthesised via reverse transcription with primers and M-Mel Reverse Transcriptase. The qPCR was carried out with an iQ5 Real-Time PCR detection system and iQ5 Optical System software (Bio-Rad). The qPCR reaction mixtures comprised 1 μL of cDNA, primer (0.6 μL), GreenMaster Mix (10 μL), and 20 μL of PCR-quality H2O.

The PCR primer sequences for ABCC11 and β-actin are shown in Table 1. Each PCR assay was done in triplicate, and each experiment included negative and no template controls, using the following thermocycle: 94°C for 10 min, then 40 cycles of 95°C (15 sec), annealing for 30 sec, and at 72°C for 25 sec. ABCC11 mRNA was quantified from qPCR data using the Pfaffl procedure, with β-actin gene as internal control [10].

| Gene  | Primer sequence                                      |
|-------|------------------------------------------------------|
| ABCC11| Sense primer (5'-CTCCACATCCTCAATTCTCTGC-3')          |
|       | Antisense primer (5'-GCCATCCATCGTGGAAGAT-3')         |
| β-actin| Sense primer (5'-GGGACGAGGCTCATCATT-3')              |
|       | Antisense primer (5'-AGCGAGCATCCCAAGTT-3')           |

Statistical analysis

Statistical significance was determined using one-way analysis of variance (ANOVA), followed by the Tukey post-hoc test (if significant). Differences were considered significant at p < 0.05. All statistical analyses were performed using GraphPad Prism (GraphPad Software, San Diego, CA, USA).

RESULTS

The MRP8 protein levels were significantly elevated in axillary tissue from untreated AO patients, when compared to healthy control tissue (p < 0.05). One month of daily “HuChou San” treatment caused a significant decrease in MRP8 protein expression in AO patient tissue, relative to tissue taken before treatment (p < 0.05). Tissues obtained after 2 and 3 months of daily “HuChou San” treatment exhibited further suppressions of MRP8 protein expression, indicating that the effect of “HuChou San” treatment depended on the duration of the treatment (Figure 1).

The ABCC11 mRNA levels were significantly elevated in axillary tissue from untreated AO patients, when compared to healthy control tissues (p < 0.01). One month of daily “HuChou San” treatment significantly down-regulated the expression of ABCC11 mRNA in AO patient tissue, relative to tissue taken before treatment (p < 0.01). Tissues obtained after 2 and 3 months of daily “HuChou San” treatment exhibited further decreases in ABCC11 mRNA levels (Figure 2).

DISCUSSION

Studies have demonstrated that human ABCC11 mRNA is produced in all foetal and adult tissues [11-13]. The ATP-binding cassette sub-family C member 11 (ABCC11) transports several biomolecules such as cyclic nucleotides, leukotriene C4 (LTC4) and S-(2,4-dinitrophenyl)-glutathione (GS-DNP);
steroid sulphates, glucuronides, and folic acid and its analogue methotrexate (MTX). The role of the MRP8 protein in the development of AO has been investigated [14,15].

The present study shows that expression levels of MRP8 protein and ABCC11 mRNA were higher in axillary skin tissue obtained from AO patients than in tissues from healthy (non-AO) individuals. In AO patients, “HuChou San” treatment time-dependently decreased the expression levels of MRP8 protein and ABCC11 mRNA in axillary skin tissue, when compared to tissues obtained from untreated patients. Abnormal expressions of MRP8 and ABCC11 may be used for clinical diagnosis of AO [9]. “HuChou San” treatment decreased MRP8 protein and ABCC11 mRNA expression levels in skin tissue from bilateral axilla of AO patients. The results of the present study provide potential new direction for the treatment of AO.

CONCLUSION

This study has shown that AO patients’ axillary skin tissue expresses markedly higher levels of ABCC11 mRNA and MRP8 protein than that of healthy patients. Treatment with “HuChou San” significantly decreases the expressions of ABCC11 mRNA and MRP8 protein in AO patients’ axillary skin tissue. Therefore, “HuChou San” may have some benefits as a therapy for AO.

DECLARATIONS

Conflict of interest

No conflict of interest is associated with this work.

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