Research Article
Serum MicroRNA Profiling and Bioinformatics of Patients with Spleen-Deficiency Syndrome

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To investigate serum microRNA (miRNA) profile and bioinformatics of patients with spleen-deficiency syndrome (SDS) and explore pathogenesis of SDS patients from miRNA levels, 10 patients with type 2 diabetes mellitus (T2DM), within which 5 patients were with SDS and the remaining were with blood stasis syndrome (BSS), and 5 healthy volunteers were recruited. Serum miRNA profiles of SDS patients were identified by quantitative PCR array. Target prediction and functional annotation for miRNAs were performed by miRSystem database. The present study identified 11 candidate serum miRNAs for SDS patients, and their targets were significantly enriched in 18 KEGG pathways and 7 GO molecular functions. Those enriched KEGG pathways included (1) metabolisms of carbohydrate, protein, amino acid, and fatty acid, (2) signaling pathways of insulin, ErbB, chemokine, calcium, and type II diabetes mellitus, (3) invasions of bacterium, Escherichia coli, and Shigella (Shigellosis), and (4) endocytosis and phagocytosis. Those enriched GO molecular functions were mainly involved in transcription regulation and regulation of metabolism. Our findings might elucidate the pathogenesis of SDS patients with disorders of substance metabolism and hypoinnunity from miRNA levels, as well as providing some miRNA biomarkers for clinical syndrome differentiation of SDS.

1. Introduction

Traditional Chinese Medicine (TCM) theory considers that spleen, also named as “Pi” in Chinese, is a comprehensive functional unit which is mainly involved in the digestive system as well as a wide variety of systems such as immune system. Spleen-deficiency syndrome (SDS) is one of the commonly encountered TCM syndromes and is closely associated with hypoinnunity and disorders of substance metabolism including nutrient digestion and absorption. Our previous studies in gene expression profile found that differentially expressed genes of SDS patients were mainly related to substance metabolism and immune regulation, and most of these genes were downregulated [1–3]. This finding had also been partially confirmed by other studies in cooperative institution [4–7]. These studies provided a new perspective for elucidating the physiological function of spleen and pathogenesis of SDS. Gene expression profiling enables researchers to observe expressions for tens of thousands of genes in a tissue simultaneously, which may be similar to the “holistic view” of TCM. However, it also has certain limitations. For example, gene expression profile from a single tissue is difficult to reflect the holistic status of multisystem disease of SDS, because of tissue specificity of gene expression. Furthermore, the differentially expressed gene is difficult to be used as biomarker for clinical diagnosis due to too many quantities.
MicroRNA (miRNA) is endogenously expressed, small single-stranded noncoding RNA molecules of 21–23 nucleotides, which regulate approximately 50% of all mammalian protein-coding genes and participate in many biological processes of cell development, differentiation, metabolism, immunity, and proliferation [8]. Serum or plasma miRNA is stable and is derived from different tissues or organs [9, 10]. Importantly, the specific serum miRNA expression profile constitutes the fingerprint of a physiological or disease condition [9]. A direct evidence from rat models showed that miRNA expression profiles in different tissues (pancreas, liver, adipose, and skeletal muscle) shared high similarity with that in blood samples [11]. Thus, serum miRNA expression profile could well reflect the holistic status of SDS patients and help to achieve comprehensive understanding of spleen essence. Moreover, serum miRNAs had been widely used in the study of biomarker and pathophysiological mechanism in different diseases, especially multisystem diseases including type 2 diabetes mellitus (T2DM) [10, 12]. However, few studies were performed in serum miRNA profiling of patients with SDS which had been recognized by TCM theory as one of the primary causes leading to the occurrence and development of T2DM.

The present study investigated serum miRNA profiling and bioinformatics of SDS patients by Exiqon serum/plasma miRNA qPCR (quantitative polymerase chain reaction) array, panel I, which is designed based on the miRNA screening results of thousands of patients with different diseases, and contains most of serum miRNAs of detectable, highly concerned and deeply studied by academic community. In addition, according to the method of combination of disease and syndrome, T2DM-BSS (Blood Stasis Syndrome) patients and HV (healthy volunteers) were selected as study control and syndrome, T2DM-BSS (Blood Stasis Syndrome) patients and HV (healthy volunteers) were selected as study control for T2DM-SDS patients to eliminate the effects of T2DM and syndrome types of TCM and to obtain the candidate serum miRNAs for SDS patients. The aim of the present study was to investigate the substance basis of miRNA and pathogenesis of SDS patients from the holistic level. It might provide further evidences for study on the scientific essence of spleen and clinical syndrome differentiation of SDS.

2. Materials and Methods

2.1. Ethics Statement. The present study was approved by the Ethics Committee of the First Affiliated Hospital of Guangzhou University of Chinese Medicine. All participants signed written informed consent.

2.2. Diagnostic Standard of T2DM. Diagnostic standard of T2DM was set up by the American Diabetes Association (ADA) as follows [13]: (1) a fasting plasma glucose (FPG) level of 126 mg/dL (7.0 mmol/L) or higher, (2) a 2-hour plasma glucose level of 200 mg/dL (11.1 mmol/L) or higher during a 75-g oral glucose tolerance test (OGTT), or (3) a random plasma glucose of 200 mg/dL (11.1 mmol/L) or higher in a patient with classic symptoms of hyperglycemia or hyperglycemic crisis.

2.3. Diagnostic Standard of Syndrome Differentiation. The diagnostic standard of syndrome differentiation was set up according to “Guiding Principle of Clinical Research on New Drugs of Chinese Medicine” and the clinical experience of authors’ institute [1, 14, 15]. Syndrome differentiation was performed by two associate chief physicians of Chinese Medicine.

The diagnostic criteria for SDS were described below. The dominant symptom items were (1) pale tongue proper with thin-white coating, (2) poor appetite, (3) abdominal distension, and (4) loose stool or diarrhea. The secondary items were (1) emaciated, (2) feebleness, and (3) weak pulse. The dominant item (1) was necessary for the diagnosis of SDS. To make the diagnosis, it also required two more dominant items or one dominant symptom plus at least two secondary symptoms.

The diagnostic criteria for BSS were as follows. The dominant symptom items were (1) localized pain, pricking, or nonpalpable abdominal pain, (2) purple-colored tongue body or tongue body with petechial, sublingual vein varicosity, (3) pathological lumps including gut swelling, neoplasm, inflammatory or noninflammatory masses, or tissue proliferation, (4) vascular ecchymosis, including cutaneous lip, gingival or acra, red thread in skin surface, or abdominal veins varicosity, (5) bleeding and melena, petechial, or bloody ascites, (6) menstrual disorders including dysmenorrhea, black or clotted menstruation, or hypogastric pain, and (7) uneven or intermittent pulse or acrotem. The secondary items were (1) scaly, dry skin, (2) limb torpidity or hemiplegia, (3) insanity, mania, amnesia, or topoparesthesia, and (4) a history of trauma, surgery, and induced abortion. The diagnosis of BSS was able to be confirmed when a patient had more than 2 dominant symptom items or 1 dominant symptom item plus at least 2 secondary symptom items.

2.4. Inclusion and Exclusion Criteria for T2DM Patients with SDS or BSS. Inclusion criteria were as follows: patients (1) were clinically diagnosed as T2DM, (2) were of the Chinese medicine syndrome type differentiated as SDS or BSS, (3) were of the age between 48 and 70 years, and (4) signed the informed consent form. Exclusion criteria were as follows: patients (1) were with other endocrine diseases in addition to T2DM, or combined with severe complications due to T2DM, (2) were with infectious or inflammatory diseases, (3) were with psychiatric or serious somatic diseases, (4) were with dyslipidemia, or (5) were in pregnancy or lactation stage.

2.5. Judgment Standard and Enrollment of Healthy Volunteers. The healthy volunteers (HV) aged between 48 and 70 years were screened by chief physician (Min Hong) through interview and physical examination. They were free of endocrine diseases including T2DM, infectious and inflammatory diseases, psychiatric and serious somatic diseases, and dyslipidemia. They should have no apparent TCM syndromes, with normal figure of tongue and pulse. They were also excluded if they were overweight or obese, presented a family history of diabetes, or long-term medication use, or failed to sign the informed consent.
2.6. Grouping. Ten patients of T2DM were recruited from the First Affiliated Hospital of Guangzhou University of Chinese Medicine during October 2013 to December 2013 and were divided into two groups according to the TCM syndrome type. Five patients (3 females and 2 males) aged from 48 to 65 (57.0 ± 7.0) years were assigned to the SDS group, while the other 5 patients (3 females and 2 males) aged from 49 to 70 (59.4 ± 9.0) years were assigned to the BSS group. Five healthy volunteers (3 females and 2 males), assigned as HV group, were also recruited from the Physical Examination Center, the First Affiliated Hospital of Guangdong Pharmaceutical University during December 2013 to April 2014. The average age of all participants was 56.4 ± 3.7 (from 51 to 61) years. No significant differences in age and sex were observed among the three study groups (P > 0.05).

2.7. Serum Sample Collection of Participants. More than 3 mL whole venous bloods were collected into a vacuum tube sans anticoagulants in a fasting state between 7:00 am and 9:00 am. After completing blood coagulation in 4 °C refrigerator for 1 hour, samples were centrifuged at 1,000 rpm for 10 min at 4 °C to obtain serum. Each 500 μL of serum was then packed in a frozen RNase-free storage tube and then stored at −80 °C until use. The whole procedure was completed within 2 hours after blood sampling.

2.8. RNA Isolation. Total RNAs, including miRNA, were isolated from the serum using Trizol reagent (Invitrogen Life Technologies, USA) according to the manufacturer’s protocol. RNA concentration and purity were determined by NanoDrop ND-1000 spectrophotometer (Thermo Fisher Scientific, USA). RNA samples that met A(absorbance)260/A280 value >1.7 and RNA concentration >60 μg/μL were used for the miRNA real-time qPCR array.

2.9. miRNA Real-Time qPCR Array. The miRCURY LNA™ universal RT miRNA PCR array, serum/plasma panel I (Exiqon, Denmark), containing 372 human mature miRNAs, was selected to detect serum miRNA expression profiles. Approximately 20–25 ng of total RNAs containing miRNA were reverse-transcribed into cDNA using the MicroRNA Reverse Transcription Kit and RT Primer Pools (Exiqon, Denmark). Then, cDNA was served as template for miRNA real-time qPCR in ABI PRISM7900 system (Applied Biosystems, USA). Thermal cycling was denatured at 95 °C for 10 min, followed by 38 cycles of 95 °C for 10 s and 60 °C for 60 s. Melting curve was performed at the end of the PCR cycle. All procedures were taken according to the manufacturer’s protocol.

2.10. Determination of Differentially Expressed miRNA. The threshold cycle (CT) values of each sample were determined by the number of PCR cycle and threshold value. Undetectable data was assigned a default CT value of 38. The CT values of individual miRNA were normalized to the housekeeping gene (SNORD38B), which had stable CT value in all samples. The miRNA expression levels for each sample were calculated using 2^\(-ΔCT\) (ΔCT = CT of target miRNA – CT of SNORD38B). Differences in miRNA expression between experimental (SDS groups) and control groups (HV or BSS group) were compared using two-tailed Student's t-test. The relative expression levels (fold changes) of miRNAs between the two groups were calculated using 2^-(ΔCT of experimental group-ΔCT of control group). The miRNAs that matched P < 0.05 and fold change >2.0 or <0.5 were defined as differentially expressed between the two groups. Data was expressed as mean ± SD. The data was analyzed with the GenEx qPCR (Exiqon, Denmark) and SPSS 18.0 (IBM, USA) software. To determine the candidate miRNAs of SDS patients, the VENN tool [16] was used to compare differentially expressed miRNAs between SDS and HV groups with that of between SDS and BSS groups via Venn diagrams.

2.11. Target Prediction of miRNA and Functional Annotation of the Predicted Target. To evaluate functions of candidate miRNAs of SDS patients, target prediction and functional annotation were performed using the miRSystem database (version 20150312) [17], which is an integrated system for characterizing enriched functions and pathways of miRNA targets. In target prediction, the miRSystem database integrates 2 experimentally validated databases, TarBase (version 7.0) and miRecords (November 2010 release), and 7 target gene prediction algorithms including Diana-microT (version 4.0), miRanda (August 2010 release), miRBridge (April 2010 release), PicTar (March 2007 release), PITA (August 2008 release), RNA22 (version 2.0), and TargetsCan (version 6.0). In functional annotation, miRSystem database integrates 5 databases including KEGG (Kyoto Encyclopedia of Genes and Genomes), Biocarta, Pathway Interaction Database (human only), Reactome (human only), and GO (Gene Ontology) molecular function. To identify significantly enriched pathways or biological function, several statistical approaches, including O/E (observed to expected) ratios of gene targets, hypergeometric and empirical P-value from permutation, and pathway ranking score from miRNA relative expression value (fold change), were also provided in the miRSystem database [17]. In the present study, miRNAs and their fold changes were submitted to miRSystem database. Only validated targets or miRNA-target interactions identified by at least 4 prediction programs were considered for annotation of KEGG pathway and GO molecular function. Moreover, parameter settings were 25–500 genes in biological functions/pathways, O/E ratios ≥1.8, and ranking score ≥2.0.

3. Results

3.1. Serum miRNA Profile of SDS Patients. Quantitative PCR arrays were introduced to compare the serum miRNA expression profiles between T2DM-SDS patients and healthy volunteers or T2DM-BSS patients. Out of 372 tested mature miRNAs, 27 miRNAs were significant differentially expressed between T2DM-SDS patients and healthy volunteers, with 5 downregulated and 22 upregulated miRNAs. Forty-three miRNAs were significant differentially expressed between patients of T2DM-SDS and T2DM-BSS, with 1 downregulated and 42 upregulated miRNAs (Figure 1). Among all these
differentially expressed miRNAs, 11 upregulated miRNAs were shared by both, which might be candidate serum miRNAs for SDS patients (Figure 1 and Table 1).

### 3.2. miRNA Target Prediction and Functional Annotation

The predicted targets of the 11 candidate serum miRNAs of SDS patients were identified by miRSystem database. Results indicated that 2,713 putative targets were regulated by these miRNAs. Among these targets, O/E ratios of 137 targets were more than 1.8. Function annotation showed that targets of these miRNAs were shared by both when P value was imported (Figure 2(a)). The 5 top canonical KEGG pathways were involved in several GO molecular functions including transcription factor activity, transcription repressor activity, RNA polymerase II transcription factor activity, and chromatin binding, and their scores were reduced in turn. The miRNAs and predicted targets in the 4 top canonical GO molecular functions were shown in Table 5.

### 4. Discussion

For the first time, the present study revealed differential serum miRNA expression profile between T2DM-SDS patients and T2DM-BSS patients or healthy volunteers by low density miRNA qPCR array. We found 11 candidate serum miRNAs of SDS patients. Target prediction and functional annotation by miRSystem database showed that targets of these miRNAs were significantly enriched in several KEGG pathways including nutrient metabolism, signaling pathway, bacterial invasion, and endocytosis and in some GO molecular functions including transcription and metabolism regulation. Our findings provided further evidences for study on the pathogenesis of SDS patients and for clinical syndrome differentiation of SDS.

TCM theory considers that the main physiological function of spleen is to govern transportation and transformation of substances. For the transportation function, spleen is responsible for digestion, absorption, and transportation of cereal essence. For the transformation function, spleen transforms cereal essence to essence, Qi (it is the basic term of TCM and related to energy), blood and body fluid in order to nourish the body. Thus spleen is closely related to the metabolism and transportation of substance including nutrients and signal transduction in Western Medicine. SDS is a TCM syndrome, which is manifested as deficiency of spleen-Qi and dysfunction of spleen governing transportation and transformation. A large number of research results in enzymology and digestive system after 1980s [18], and a
Table 2: KEGG pathways from 11 candidate serum miRNAs of SDS patients.

| Term                               | Term ID | Targets in the term | miRNAs in the term | P value  |
|------------------------------------|---------|---------------------|--------------------|----------|
| Insulin signaling pathway          | hsa04910| 4                   | 8                  | 0.0090   |
| ErbB signaling pathway             | hsa04012| 3                   | 7                  | 0.0150   |
| Chemokine signaling pathway        | hsa04062| 4                   | 7                  | 0.0240   |
| Endocytosis                        | hsa04144| 4                   | 6                  | 0.0286   |
| Calcium signaling pathway          | hsa04020| 4                   | 5                  | 0.0198   |
| Carbohydrate digestion and absorption | hsa04973| 2                   | 5                  | 0.0270   |
| Ubiquitin mediated proteolysis     | hsa04120| 3                   | 5                  | 0.0421   |
| Bacterial invasion of epithelial cells | hsa05100| 4                   | 4                  | 0.0009   |
| FcγR-mediated phagocytosis          | hsa04666| 3                   | 4                  | 0.0181   |
| Arginine and proline metabolism    | hsa00330| 2                   | 4                  | 0.0400   |
| Galactose metabolism               | hsa00052| 2                   | 3                  | 0.0108   |
| Amino sugar and nucleotide sugar metabolism | hsa00520| 2                   | 3                  | 0.0315   |
| Starch and sucrose metabolism      | hsa00500| 2                   | 3                  | 0.0387   |
| Pathogenic *Escherichia coli* infection | hsa05130| 3                   | 2                  | 0.0047   |
| Valine, leucine, and isoleucine degradation | hsa00280| 2                   | 2                  | 0.0281   |
| Type II diabetes mellitus          | hsa04930| 2                   | 2                  | 0.0315   |
| Shigellosis                        | hsa05131| 2                   | 2                  | 0.0489   |
| Fatty acid metabolism              | hsa00071| 3                   | 1                  | 0.0023   |

Table 3: GO molecular functions from 11 candidate serum miRNAs of SDS patients.

| Term                               | Term ID | Targets in the term | miRNAs in the term | P value  |
|------------------------------------|---------|---------------------|--------------------|----------|
| Transcription repressor activity   | GO:0016564| 6                   | 9                  | 0.0069   |
| Enzyme activator activity          | GO:0008047| 6                   | 9                  | 0.0148   |
| Lipid binding                      | GO:0008289| 6                   | 9                  | 0.0222   |
| Transcription factor binding       | GO:000989| 7                   | 7                  | 0.0061   |
| Transcription factor activity      | GO:003406| 4                   | 5                  | 0.0024   |
| Carboxylic acid binding            | GO:003682| 4                   | 5                  | 0.0240   |
| Chromatin binding                  | GO:003702| 5                   | 4                  | 0.0186   |

Figure 2: Venn diagrams of KEGG pathway and GO molecular function annotations based on P and score value. Score 1 indicated number of enriched biological functions in ranking score > 2.0 from T2DM-SDS patients versus HV. Score 2 indicated number of enriched biological functions in ranking score >2.0 from T2DM-SDS patients versus T2DM-BSS patients. (a) denoted KEGG pathway and (b) denoted GO molecular function.
Table 4: miRNAs and their targets in 5 top canonical KEGG pathways (score down).

| Term                      | Targets in the term                          | miRNAs in the term         |
|---------------------------|-----------------------------------------------|-----------------------------|
| **ErbB signaling pathway**| ELK1: ELK1, member of ETS oncogene family     | miR-135a-3p, miR-135b-5p, miR-136-3p, miR-219a-5p |
|                           | SHC1: SHC (Src homology 2 domain containing) transforming protein 1 | miR-124-3p, miR-219a-5p, miR-9-5p, miR-96-5p |
|                           | SHC3: SHC (Src homology 2 domain containing) transforming protein 3 | miR-124-3p |
|                           | ELK1                                          | As above                   |
|                           | SHC1                                          | As above                   |
| **Insulin signaling pathway** | HK3: hexokinase 3 (white cell)                | miR-137                     |
|                           | SHC3                                          | As above                   |
|                           | GNAQ: guanine nucleotide binding protein (G protein), q polypeptide | miR-135a-5p, miR-135b-5p, miR-96-5p |
|                           | SLC25A5: solute carrier family 25 (mitochondrial carrier; adenine nucleotide translocator), member 5 | miR-135a-5p, miR-135b-5p, miR-137 |
| **Calcium signaling pathway** | CACNA1G: calcium channel, voltage-dependent, T type, alpha 1G subunit | miR-137, miR-96-5p |
|                           | SPHK2: sphingosine kinase 2                   | miR-137, miR-153-3p         |
|                           | GRK6/G protein-coupled receptor kinase 6      | miR-137, miR-141-3p, miR-200a-3p |
|                           | EPNI: epsin 1                                 | miR-141-3p, miR-200a-3p     |
| **Endocytosis**           | STAMBBP: STAM binding protein                 | miR-135a-5p, miR-135b-5p    |
|                           | CHMP4A: charged multivesicular body protein 4A | miR-96-5p                 |
|                           | SHC1                                          | As above                   |
| **Bacterial invasion of epithelial cells** | ARPC1A: actin related protein 2/3 complex, subunit 1A, 41 kDa | miR-9-5p |
|                           | ARPC1B: actin related protein 2/3 complex, subunit 1B, 41 kDa | miR-124-3p |
|                           | SHC3                                          | As above                   |

few results in gene expression profiles after 2000s [1, 2, 5, 6], had confirmed that SDS patients had the disorder of digestion and absorption and the abnormal metabolism of substances. Nonetheless, the understanding of the pathogenesis of SDS was not comprehensive because of particularity of SDS disease itself and shortages of these research methods. Spleen was a comprehensive functional unit and its anatomical location is not clear; thus pathogenesis of SDS was complex and complicated. One or several indicators were difficult to fully reflect the pathogenesis of SDS. Moreover, gene expression profiling was not suitable for study of multisystem diseases. However, serum miRNA expression profile, as a novel method, could overcome these shortcomings as mentioned in the introduction. Moreover, serum miRNA expression profile in SDS patients had not been reported.

In the present study, for the first time, we reported 11 serum miRNAs from SDS patients and all these miRNAs were upregulated. Among these 11 miRNAs, miR-9, miR-96, miR-153, and miR-200a had been confirmed to be related to the occurrence of type 2 diabetes [10, 12, 19]. Furthermore, miR-9, miR-96, and miR-153 regulated insulin secretory [12, 19], while miR-200a and miR-96 regulated insulin receptor pathway [12]. Besides, miR-124-3p controlled glucagon release by directly targeting iGluR2 and iGluR3 [20]; miR-137 regulated proliferation and differentiation of human adipose tissue stromal cells by targeting CDC42 [21]; miR-135a-5p and miR-135b-5p controlled adipogenesis and atherosclerosis, respectively [22, 23]; miR-141-3p contributed to mitochondrial dysfunction in high-fat-diet induced obesity by inhibiting PTEN [24]. These functional studies in miRNAs related to SDS patients showed that there were abnormal metabolism of glucose and lipid in SDS patients from the macroscopic view. In addition, functional annotation in the present study showed that targets downregulated by 11 miRNAs of SDS patients were significantly enriched in several KEGG pathways related to nutrient metabolism, signaling pathway, and endocytosis and in some GO molecular functions including regulation of transcription and metabolism. These findings further indicated SDS patients had dysfunction of nutrient metabolism and transportation and signal transduction. Duan et al. found that SDS model rats had lower expressions of key factors from Ca^2+/CaMK II signaling pathways in...
Table 5: miRNAs and their targets in 4 top canonical GO molecular functions (score down).

| Term | Targets in the term | miRNAs in the term |
|------|---------------------|--------------------|
| Transcription factor binding transcription factor activity | TAIF2: TAIF2 RNA polymerase II, TATA box binding protein (TBP) associated factor, 20 kDa | miR-137, miR-141-3p, miR-200a-3p |
| | CSDA | As above |
| | DYRK1B: dual-specificity tyrosine-(Y)-phosphorylation regulated kinase 1B | miR-135a-5p, miR-135b-5p, miR-9-5p |
| | Pias2: protein inhibitor of activated STAT 2 | miR-124-5p, miR-135a-5p, miR-135b-5p |
| | DDX5: DEAD (Asp-Glu-Ala-Asp) box helicase 5 | As above |
| | SIRT1: sirtuin 1 | miR-124-3p, miR-135a-5p, miR-135b-5p |
| | ELK3: ELK3, ETS-domain protein (SRF accessory protein 2) | miR-124-3p, miR-135a-5p, miR-135b-5p |
| | TWIST2: twist family bHLH transcription factor 2 | miR-124-3p |
| | REST: REI-silencing transcription factor | miR-9-5p |
| | NKAP: NFkB activating protein | miR-124-3p |
| | YBX1: Y box binding protein 1 | miR-153-3p, miR-135a-5p, miR-135b-5p, miR-219a-5p |
| | KLF4: Kruppel-like factor 4 (gut) | As above |
| | SIRT1 | As above |
| | TAIF2 | As above |
| | CSDA | As above |
| RNA polymerase II transcription factor activity | HTATSF1: HIV-1 Tat specific factor 1 | miR-141-3p |
| | MNX1: motor neuron and pancreas homeobox 1 | miR-200a-3p, miR-141-3p |
| | PIAS2 | As above |
| | REST | As above |
| | NKAP | As above |
| Chromatin binding | ING2: inhibitor of growth family, member 2 | miR-153-3p |
| | YBX2: Y box binding protein 2 | miR-135a-5p, miR-135b-5p |

skeletal muscle than control rats [25]. In our previous studies in chronic superficial gastritis patients (CSG) with SDS [1, 2], we found that more than 70% differentially expressed genes between CSG-SDS patients and CSG-SSHS (spleen-stomach hygropyrexia syndrome) patients or healthy volunteers were associated with metabolism of nutrients including carbohydrate, protein, amino acid, lipid and nucleic acid, and metal ion metabolism. Furthermore, genes related to carbohydrate metabolisms participated in glycogen and glycogen metabolic processes; genes related to protein metabolism participated in signaling pathways, amino acid metabolism, and protein ubiquitination and targeting transport including endocytosis; genes related to nucleic acid metabolism participated in regulations of replication and transcription; genes related to metal ion metabolism were mainly calcium binding protein. Interestingly, among all these genes involved in metabolism, more than 80% genes were downregulated in GSG-SDS patients. Moreover, downregulated ACADVL (acyl-CoA dehydrogenase, very long chain) and ACAA2 (acyl-coenzyme A acyltransferase 2) [1, 2, 5], which catalyze the first and last steps of the mitochondrial fatty acid beta-oxidation pathway, were targeted by upregulated 135a-5p and miR-124-3p in current study, respectively. Thus, our previous reports of gene expression profiles from CSG-SDS patients strongly supported the current research results. They elucidated scientific connotation of spleen governing transportation and transformation and pathogenesis of SDS patients with disorders of substance metabolism from gene and miRNA levels, respectively. In addition, in the present study, we found that all the KEGG pathways and GO molecular functions related to nutrient metabolism were deleted from the significantly enriched biological functions when weighted pathway-ranking score \( \geq 2.0 \) was used to identify the enriched biological functions. These findings
might illustrate that spleen governing transportation was more important than spleen governing transformation in physiological function of spleen.

TCM theory considers that spleen is the foundation of acquired constitution and that the reinforced function of spleen is responsible for individual's insusceptibility to pathogenic factors in the four seasons. Hence, spleen can protect the body from external pathogenic factor, and thus spleen-deficiency would break out all kinds of diseases and ailments. Accumulating clinical observations has indicated that SDS patients usually manifest weak constitution and hypoimmunity, as well as dysfunction of the digestive system. Luo et al. found that the pathogenesis of SDS patients had genomic basis associated with immune, and SDS patients had disorder of the immune functions [7]. In the present study, we found that targets downregulated by the 11 miRNAs of SDS patients were significantly enriched in 3 KEGG pathways related to invasion of pathogenic factor, including bacterium, Escherichia coli, and Shigella (Shigellosis). Furthermore, KEGG pathway of bacterial invasion of epithelial cells was one of the 5 top canonical KEGG pathways. These findings strongly indicated that SDS patients had a reduced resistance to external pathogenic factor. In our recent studies about gene expression profile from peripheral white blood cells of T2DM-SDS patients [3, 15], we found that approximate 30% of differentially expressed genes between T2DM-SDS patients and T2DM-BSS patients or healthy volunteers were associated with immune regulation or response. Furthermore, more than 60% genes were downregulated in all these differentially expressed genes. Our previous studies of gene expression profiles from T2DM-SDS patients supported the current study. In addition, among 11 miRNAs of SDS patients in the present study, miR-219-5p was confirmed to directly regulate inflammatory adipokine expression and macrophage innate immune response [26]. Hence, all these results elucidated scientific connotation of spleen governing protection of the body from external pathogenic factor and pathogenesis of SDS patients with hypoimmunity from gene and miRNA levels, respectively.

Although the present study obtained some meaningful results in the scientific essence of spleen and pathogenesis of SDS patients, there were some possible limitations. For example, the present sample size was limited, and verifications of predicted targets were not performed. Thus further researches were needed in the future.

Disclosure
All of the funders had no further role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

Competing Interests
The authors declared that there was no conflict of interests regarding the publication of this paper.

Authors’ Contributions
Ze-Min Yang and Long-Hui Chen wrote the manuscript; Ze-Min Yang and Wei-Wen Chen were the principal investigators and together contributed to the development of the overall research plan, study protocol and oversight, and analyzed the data; Min Hong, Ying-Yu Chen, Long-Hui Chen, and Xiao-Rong Yang were responsible for participant recruitment and sample collection. Si-Meng Tang, Qian-Fa Yuan, and Zhen-Yu He performed the miRNA qPCR array experiments. All authors revised the manuscript and approved the final version.

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