Research Article

Fufang Xueshuuantong for Diabetic Kidney Disease: A Systematic Review and Meta-Analysis

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Objective. The objective of this meta-analysis was to systematically assess the efficacy and safety of patented Chinese medicine Fufang Xueshuuantong (FFXST) for the treatment of diabetic kidney disease (DKD).

Methods. Randomized controlled trials (RCTs) of FFXST for DKD treatment were searched until May 31, 2020, in seven electronic databases: PubMed, Embase, Cochrane Library, CNKI, Wanfang, VIP, and Chinese Biomedical Literature. The Cochrane risk test from the Cochrane Handbook was used as a bias tool to assess the methodological quality, and Review Manager (RevMan) 5.3 was used to analyze the results. Grading of Recommendations Assessment, Development, and Evaluation (GRADE) criteria were used to classify the quality of evidence.

Results. Thirteen RCTs involving 1,186 patients were included. The meta-analysis revealed that the efficacy of FFXST in treatment of DKD was significantly superior to that of the control treatment ($P<0.0006$). The urinary albumin excretion rate ($P<0.01$), urinary albumin creatinine ratio ($P<0.0001$), and microalbumin ($P<0.0001$) were lower in the treatment groups than in the control group. There was also a decrease in low-density lipoprotein cholesterol ($P<0.001$), serum triglyceride ($P<0.001$), and C-reactive protein ($P<0.0001$) in the treatment groups compared with those in the control group. No significant difference in hemoglobin A1c level ($P=0.76$) and systolic blood pressure ($P=0.34$) was noted between the treatment and control groups. Three studies reported adverse events, including dizziness and intolerance. In the other 10 trials, adverse events were not mentioned. Conclusion. FFXST appears to be effective in the treatment of DKD. However, the low methodological quality of the RCTs suggests that larger, better-designed RCTs are required to verify the clinical effectiveness and safety of FFXST.

1. Introduction

Diabetic kidney disease (DKD) is one of the most important microvascular complications of diabetes, as well as a key cause of end-stage renal disease (ESRD). It also increases the risk of cardiovascular disease and all-cause death in patients with diabetes [1, 2]. With the incidence of diabetes increasing annually, the number of DKD cases is also increasing. Approximately 20%–40% of patients with diabetes also have DKD [3]. The risk factors of DKD include age, disease course, blood pressure, obesity (especially abdominal obesity), blood lipid, uric acid, and environmental pollutants [4]. The main clinical manifestations of DKD are proteinuria and (or) impaired glomerular filtration rate (GFR) [3, 5].
to develop additional treatment methods to counter DKD [6].

In traditional Chinese medicine, DKD belongs to the categories “cloudy urine” and “edema” [7]. In recent years, many Chinese herbal extracts and Chinese patent medicines have demonstrated the reduction of proteinuria and the improvement of renal function in the treatment of DKD. Among them, the Chinese patent medicine Fufang Xueshuantong (FFXST) has been widely used in the treatment of DKD and other diabetic microvascular complications in China. FFXST is composed of Notoginseng Radix, Radix Astragali, Salvia Miltiorrhiza, and Scrophularia Ningpoensis. Studies [8–10] have shown that FFXST has a protective effect on the kidneys of diabetic rats, can reduce oxidative stress injury, regulate the RASS system, promote podocyte repair, and improve microcirculation and antiplatelet aggregation.

Although several clinical trials have suggested the efficacy of FFXST for DKD, most of the trials have been single-center, including small cohorts and highly different treatment schemes. Hence, it is difficult to verify the clinical efficacy of these treatment strategies. Therefore, the goal of this meta-analysis was to assess the efficacy and safety of FFXST for the treatment of DKD, providing evidence for clinical practice.

2. Materials and Methods

2.1. Search Strategy. We searched seven electronic databases, including PubMed database, Embase database, Cochrane Library, China National Knowledge Infrastructure database, Wanfang Database, Chinese Science Journal Database, and the Chinese Biomedical Database. We retrieved studies from all of these databases published before May 31, 2020. Our search keywords were as follows: “diabetic kidney disease” OR “diabetic nephropathy” AND “fufang xueshuantong” OR “compound xueshuantong” AND “randomized controlled trial,” “controlled clinical trial,” “random,” “randomly,” “randomized” OR “control.” Furthermore, we manually searched additional relevant publications according to reference lists from the resulting publications. Different search strategies were applied to Chinese and foreign language databases, without restriction on language or publications.

2.2. Inclusion Criteria

(1) Types of trials: randomized controlled trials (RCTs) using FFXST monotherapy or combination therapy with western medicine for the treatment of DKD were included.

(2) Types of patients: regardless of the type of diabetes mellitus (DM), stage of the DKD (Mogensen staging criteria), age, gender, or race, we recruited patients who were diagnosed with DKD by clearly defined or internationally recognized criteria.

(3) Types of interventions: the experimental group was treated with FFXST monotherapy irrespective of the dosage form or combined with conventional western medicine (ACEI/ARB). There was no limit to interventions in control groups, whether placebo or ACEI/ARB. Additionally, both groups received routine treatment, such as treatment to lower blood pressure, controlling blood glucose, and regulating serum lipids.

(4) Types of outcomes: all included studies that reported at least one of the following outcomes: total effective rate or proteinuria indicators.

2.3. Exclusion Criteria

(1) Interventions that included other traditional Chinese medicine (TCM) therapies, such as Chinese patent medicine, TCM decoction, herbal extracts, or acupuncture were excluded.

(2) Studies with erroneous or incomplete data were excluded.

(3) Duplicate publications were excluded.

2.4. Data Extraction. Two researchers extracted the information independently. The data included study ID, baseline patients, disease data, interventions, and outcomes (e.g., sample size, age, gender, type of DM, stage of DKD, interventional measures, treatment duration, reporting of adverse events, and outcome measures). Discrepancies were resolved by discussion with other authors.

2.5. Quality Assessment. Two researchers assessed the risk of bias in trials based on the Cochrane Handbook for the methodological quality of the included studies. We applied the RevMan5.3 to assess the following six items: random sequence generation (selection bias), allocation concealment (selection bias), blinding of participants and personnel (performance bias), blinding of outcome assessment (detection bias), incomplete outcome data (attrition bias), selective reporting (reporting bias), and other sources of bias such as baseline comparability of subjects and sample size. We also used the Grading of Recommendations, Assessment, Development, and Evaluations (GRADE) approach to assess the quality of evidence for each outcome by GRADEpro GDT software. This classifies evidence as high, moderate, low, or very low quality. Discrepancies were resolved by a third party (Qing Ni).

2.6. Data Analysis. RevMan 5.3 software (Cochrane Collaboration, Oxford, UK) was applied for statistical analysis. Dichotomous data were presented as relative risk (RR), and continuous data were included as the mean difference (MD) or standardized mean difference (SMD) and both included a 95% confidence interval (CI). The heterogeneity evaluations were conducted using a Chi² test. The fixed-effects model was used when the heterogeneity was significant (P > 0.10, I² ≤ 50%); otherwise, a randomized effects model was used (i.e., when P < 0.10, I² ≥ 50%). The possible sources of heterogeneity were explored by sensitivity analysis and
subgroup analysis. Publication bias was tested using funnel plots when the number of experiments was ≥10 [11].

2.7. Outcomes. The primary outcome indicator was a total effective rate, which was based on changes in symptoms and the level of proteinuria [12]. The total effective rate was categorized as significantly effective cases (urinary albumin excretion rate [UAER] returned to normal levels or decreased by more than 50%, with an obvious improvement in symptoms), effective cases (UAER decreased by less than 50%, improvement in symptoms), or ineffective cases (no improvement in either UAER and symptoms).

The secondary outcomes included the proteinuria indicators UAER, urinary albumin creatinine ratio (ACR), and microalbumin (mAlb); the renal function indicators estimated glomerular filtration rate (eGFR), blood urea nitrogen (BUN), and serum creatinine (Scr); hemoglobin A1c (HbA1c); low-density lipoprotein cholesterol (LDL-C); triglyceride (TG) level; blood pressure indicators; inflammatory indicators.

3. Results

3.1. Study Search and Selection. Initially, a total of 114 publications were identified from the seven electronic databases. After removing 67 duplicate publications, we excluded 20 nonclinical studies by reading the titles and abstracts. After a full-text review, we excluded two studies with significant data errors, three nonrandomized controlled studies, and nine interventions or outcome indicators that did not meet inclusion criteria. Finally, 13 studies were included in this meta-analysis. The retrieval process is shown in Figure 1.

3.2. Characteristics of the Included Studies. Characteristics of the 13 studies [13–25] are summarized in Table 1. All the included studies were published between 2009 and 2019. The 13 RCTs involved 1186 subjects (592 in treatment groups and 594 in control groups), and the sample size for each study ranged from 48 to 130 subjects. In terms of the disease type and stage, patients with type 2 diabetes (T2DM) were included in eight studies, whereas the remaining five studies did not report in detail the type of diabetes patients included. Except for one study [14] that did not report the DKD stage, the remaining 12 studies included subjects who were DKD patients in Mogensen III according to the Mogensen stage. The subject's DKD diagnosis was clear in 13 studies. All studies included patients who met the diagnostic criteria for DKD, nine of which used the criteria of the World Health Organization (WHO) DM diagnostic criteria [26] and Mogensen diagnostic [27]. One study [18] referred to the American Diabetes Association criteria combined with the Epidemiology and Diagnostic Criteria of Diabetic Nephropathy [28]. Another [16] used the WHO DM1999 and pathological diagnosis of renal biopsy. The diagnosis of DKD in one study [14] was based on internal diagnostic criteria for diabetes combined with symptoms of proteinuria and history of diabetic retinopathy and the internal DKD diagnostic criteria were used in another study [24].

Compared with the control group, treatment groups in six RCTs [16, 17, 22–25] were treated with FFXST monotherapy, whereas treatment groups in the other seven RCTs received FFXST combined with ACEI/ARB. All patients in both groups were treated with conventional hypoglycemic therapy. The duration of the trials ranged from 8 weeks to 24 weeks.

Only two studies used the total effective rate based on changes in symptoms and proteinuria levels as the main outcome indicators. In terms of proteinuria indicators, five studies reported UAER, six studies reported ACR, and six studies reported mAlb. Additionally, we also used BUN, HbA1c, standard bicarbonate (SBP), TG, LDL-C, and C-reactive protein (CRP) as secondary outcome indicators. Adverse events were not mentioned in three studies [15, 17, 21], and none of the studies reported a decrease in the quality of life or took adverse indicators (e.g., deterioration rate, access to dialysis rate, etc.) as outcome measures.

3.3. Risk of Bias in the Included Studies. The methodological quality assessment of the 13 RCTs is shown in Figures 2 and 3. Two studies [16, 17] adopted methods of randomization using a random number table; one study [20] used mechanical random sampling, and seven studies [13, 14, 18, 19, 22, 23, 25] only mentioned “randomization” but did not describe specific methods. The remaining three RCTs had a high risk of selection bias, because two [15, 21] followed the order of medical treatment, and one [24] followed the case number. None of the 13 studies were double-blind, and no study indicated details on allocation concealment or sample size calculations. Two studies [22, 23] showed high-risk bias in selective reporting. Baseline information was similar for different groups of subjects in all 13 studies. In short, the quality of all RCTs was generally low and contained a risk of bias.

3.4. Effects of the Interventions

3.4.1. Total Effective Rate. The total effective rate was reported in two studies, and the results indicated significant differences between the two groups. These trials exhibited nonsignificant heterogeneity ($\chi^2 = 0.34$, $P = 0.56$, I2 = 0%); thus, the fixed-effects model was used for statistical analysis. The total effective rate of the treatment groups was superior to that of the control groups ($N = 177$, RR = 1.37, 95% CI: 1.15–1.64, $Z = 3.45$, $P = 0.0006$) (Figure 4).

3.4.2. UAER. Five studies (Dai XM, 2012, Wang ML, 2017, Wang NN, 2012, Yang P, 2014, and Yun P, 2013) evaluated changes in UAER (Figure 5) according to indicators involving 383 patients (MD = −30.98, 95% CI: −49.30–12.66, $Z = 3.31$, $P = 0.0009$). There was significant heterogeneity among the studies ($\chi^2 = 36.88$, $P < 0.00001$, I2 = 89%), and a random effects model was used for combined analysis.
To further compare the differences in UAER between FFXST combined with conventional western medicine and control groups, subgroup analysis was performed. In one study [17], FFXST monotherapy in the treatment group was compared with that of the control group (MD $= -51.60$, 95% CI: $-73.00$–$39.16$, $Z = 8.13$, $P < 0.00001$). FFXST combined with ACEI/ARB was compared with control groups in the remaining four studies ($N = 305$, MD $= -24.61$, 95% CI: $-57.00$–$9.02$, $Z = 3.09$, $P = 0.002$), without significant heterogeneity ($\chi^2 = 13.44$, $P = 0.004$, $I^2 = 78\%$). Sensitivity analysis suggested that the heterogeneity between the subgroups decreased significantly after removal of one study [21] ($\chi^2 = 0.66$, $P = 0.72$, $I^2 = 0\%$). The test method and kit used in this study for the detection of UAER indicators may have been a source of heterogeneity.

3.4.3. ACR. Six studies involving a total of 560 patients reported ACR as an outcome (SMD $= -1.33$, 95% CI: $-1.90$–$0.76$, $Z = 4.55$, $P < 0.00001$) (Figure 6). A random effects model was used because of the significant heterogeneity among studies ($\chi^2 = 53.72$, $P < 0.00001$, $I^2 = 91\%$). Among these, two studies compared the ACR outcomes of FFXST plus ACEI/ARB with those of the control group ($N = 250$, SMD $= -0.6$, 95% CI: $-0.85$–$0.34$, $Z = 4.62$, $P < 0.00001$) without heterogeneity ($\chi^2 = 0.35$, $P = 0.55$, $I^2 = 0\%$). Four studies compared ACR outcomes between FFXST alone and control groups ($N = 310$, SMD $= -1.71$, 95% CI: $-2.22$–$-1.20$, $Z = 6.55$, $P < 0.00001$), with heterogeneity ($\chi^2 = 14.06$, $P = 0.003$, $I^2 = 79\%$). Further sensitivity analysis showed that, after the exclusion of one study [17],
<table>
<thead>
<tr>
<th>No.</th>
<th>Study ID</th>
<th>Sample size (T/C)</th>
<th>Average age (T)</th>
<th>Average age (C)</th>
<th>Sex (M/F)</th>
<th>Type of DM</th>
<th>DM duration (year)</th>
<th>Stage of DKD</th>
<th>Diagnostic criteria</th>
<th>Intervention (T)</th>
<th>Intervention (C)</th>
<th>Treatment duration (weeks)</th>
<th>Treatment event report</th>
<th>Outcomes</th>
</tr>
</thead>
<tbody>
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<td>1</td>
<td>Dai XM [13]</td>
<td>40/40</td>
<td>51.7 ± 5.2</td>
<td>52.2 ± 1.3</td>
<td>T: 25/15 C: 23/17</td>
<td>No restriction</td>
<td>T: 12 ± 6.1 C: 13 ± 5.9</td>
<td>Mogensen III</td>
<td>WHO DM1999 +MDC</td>
<td>FFXST (1.5 g. tid) + ACEI (Benazepril 10 mg, qd)</td>
<td>ACEI (Benazepril 10 mg, qd)</td>
<td>12 w</td>
<td>NO</td>
<td>Total effective rate, UAER, TG</td>
</tr>
<tr>
<td>2</td>
<td>Li T [14]</td>
<td>65/65</td>
<td>46.5 ± 1.38</td>
<td>48.2 ± 1.0</td>
<td>T: 4/24 C: 37/28</td>
<td>T2DM</td>
<td>T: ≥5 C: ≥5</td>
<td>Not report</td>
<td>Internal diagnostic criteria for diabetes combined with symptoms of proteinuria and history of diabetic retinopathy</td>
<td>FFXST (1.5 g. tid) + ACEI/ARB</td>
<td>FFXST (1.5 g. tid) + ACEI/ARB</td>
<td>12 w</td>
<td>NO</td>
<td>ACR, TG, LDL-C</td>
</tr>
<tr>
<td>3</td>
<td>Lu YZ [15]</td>
<td>29/29</td>
<td>48.7 ± 4.26</td>
<td>50.37 ± 4.8</td>
<td>T: 15/15 C: 14/16</td>
<td>T2DM</td>
<td>T: 5.78 ± 0.15 C: 5.85 ± 0.08</td>
<td>Mogensen III</td>
<td>WHO DM1999 + MDC</td>
<td>FFXST (1.5 g. tid) + ACEI</td>
<td>ACEI (Benazepril 10–20 mg, qd)</td>
<td>12 w</td>
<td>YES</td>
<td>mAlb, HbA1c, TG, SBP, CRP</td>
</tr>
<tr>
<td>4</td>
<td>Peng SL 2015 [16]</td>
<td>60/60</td>
<td>59.89 ± 8.24</td>
<td>58.28 ± 8.5</td>
<td>T: 34/28 C: 26/26</td>
<td>T2DM</td>
<td>T: 7.2 ± 3.1 C: 7.5 ± 2.9</td>
<td>Mogensen III</td>
<td>WHO DM1999 + pathological diagnosis of renal biopsy</td>
<td>FFXST (1.5 g. tid) + ARB (Valsartan 10–20 mg, qd)</td>
<td>ALB (Valsartan 80 mg, qd)</td>
<td>8 w</td>
<td>NO</td>
<td>ACR, mAlb, BUN, TG</td>
</tr>
<tr>
<td>5</td>
<td>Wang ML 2017 [17]</td>
<td>40/38</td>
<td>54.5 ± 6.6</td>
<td>56.1 ± 7.1</td>
<td>T: 22/18 C: 19/19</td>
<td>T2DM</td>
<td>—</td>
<td>Mogensen III</td>
<td>WHO DM1999 + MDC</td>
<td>FFXST (1.5 g. tid) + ARB (Losartan 50 mg, qd)</td>
<td>ARB (Losartan 50 mg, qd)</td>
<td>24 w</td>
<td>NO</td>
<td>UACR, BUN, HbA1c, TG, LDL-C</td>
</tr>
<tr>
<td>6</td>
<td>Wang ML [18]</td>
<td>38/42</td>
<td>55.4 ± 5.4</td>
<td>54.5</td>
<td>T: 17/21 C: 18/24</td>
<td>T2DM</td>
<td>—</td>
<td>Mogensen III</td>
<td>ADA2009 + EDCDN</td>
<td>FFXST (1.5 g. tid) + ARB (Losartan 50 mg, qd)</td>
<td>ARB (Losartan 80 mg, qd)</td>
<td>12 w</td>
<td>NO</td>
<td>ACR, HbA1c, TG, SBP</td>
</tr>
<tr>
<td>7</td>
<td>Wu NN 2012 [19]</td>
<td>60/60</td>
<td>56.3 ± 11.5</td>
<td>58.7 ± 10.2</td>
<td>T: 35/25 C: 34/26</td>
<td>T2DM</td>
<td>—</td>
<td>Mogensen III</td>
<td>WHO DM1999 + MDC</td>
<td>FFXST (1.5 g. tid) + ACEI</td>
<td>ACEI (Benazepril 10 mg, qd)</td>
<td>12 w</td>
<td>NA</td>
<td>UACR, BUN, HbA1c, TG, LEL-C</td>
</tr>
<tr>
<td>8</td>
<td>Yang P 2014 [20]</td>
<td>24/24</td>
<td>52.1 ± 5.3</td>
<td>53.6 ± 4.9</td>
<td>T: 13/11 C: 12/12</td>
<td>T2DM</td>
<td>—</td>
<td>Mogensen III</td>
<td>WHO DM1999 + MDC</td>
<td>FFXST (1.5 g. tid) + ARB (Valsartan 80 mg, qd)</td>
<td>ARB (Valsartan 80 mg, qd)</td>
<td>12 w</td>
<td>NO</td>
<td>ACR, HbA1c, TG, SBP</td>
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<tr>
<td>9</td>
<td>Yun P 2013 [21]</td>
<td>51/51</td>
<td>53.5 ± 6.4</td>
<td>55.1 ± 7.2</td>
<td>T: 32/19 C: 29/22</td>
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<td>—</td>
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<td>WHO DM1999 + MDC</td>
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<td>ARB (Losartan 50 mg, qd)</td>
<td>12 w</td>
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<td>Total effective rate, UACR, BUN, HbA1c, TG, LEL-C, SBP</td>
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<td>10</td>
<td>Zhang JH [22]</td>
<td>45/45</td>
<td>—</td>
<td>—</td>
<td>50/40</td>
<td>No restriction</td>
<td>8.5</td>
<td>Mogensen III</td>
<td>WHO DM1999 + MDC</td>
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<td>ARB (Losartan 50 mg, qd)</td>
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<td>NO</td>
<td>mAlb, BUN</td>
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<td>11</td>
<td>Zhang Z [23]</td>
<td>45/45</td>
<td>—</td>
<td>—</td>
<td>50/40</td>
<td>No restriction</td>
<td>8.5</td>
<td>Mogensen III</td>
<td>WHO DM1999 + MDC</td>
<td>FFXST (1.5 g. tid) + ARB (Losartan 50 mg, qd)</td>
<td>ARB (Losartan 50 mg, qd)</td>
<td>12 w</td>
<td>NO</td>
<td>mAlb, CRP</td>
</tr>
<tr>
<td>12</td>
<td>Zhang Z [24]</td>
<td>45/45</td>
<td>48.9 ± 10.1</td>
<td>49.4 ± 9.6</td>
<td>T: 31/14 C: 29/16</td>
<td>No restriction</td>
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<td>The internal DKD diagnostic criteria</td>
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<td>12 w</td>
<td>NO</td>
<td>ACR, mAlb, HbA1c</td>
</tr>
<tr>
<td>13</td>
<td>Zhang Z [25]</td>
<td>50/50</td>
<td>58.9 ± 10.18</td>
<td>59.2 ± 9.7</td>
<td>60/40</td>
<td>No restriction</td>
<td>—</td>
<td>Mogensen III</td>
<td>The internal DKD diagnostic criteria</td>
<td>FFXST (1.5 g. tid)</td>
<td>Blank</td>
<td>12 w</td>
<td>NO</td>
<td>ACR, mAlb, HbA1c</td>
</tr>
</tbody>
</table>

the subgroup sensitivity decreased ($\chi^2 = 14.06, P = 0.032, I^2 = 12\%$). The sources of heterogeneity may have been related to the conventional treatment regimen adopted in the study, where ACEI/ARB drugs were preferred for patients with hypertension, leading to some patients belonging to the FFXST plus ACEI/ARB subgroup, rather than FFXST subgroup alone.

### mAlb

Six studies reported mAlb as an outcome ($N = 548$, MD $= -36.29$, 95% CI: $-54.45$–$-18.13, Z = 3.92, P < 0.0001$) (Figure 7) with heterogeneity ($\chi^2 = 596.01, P < 0.00001, I^2 = 99\%$) and the use of a random effects model. Then, we performed a subgroup analysis to compare the effects of different diabetes types on the results. The subjects in two studies were T2DM patients, and there was no heterogeneity between these groups ($\chi^2 = 0.00, P = 0.97, I^2 = 0\%$). The results showed that the mAlb changes were statistically different between treatment and control groups ($N = 178$, MD $= -22.32$, 95% CI: $-27.56$–$-17.09, Z = 8.36, P < 0.00001$). The other four studies did not limit the type of diabetes and had significant heterogeneity ($\chi^2 = 564.54, P < 0.00001, I^2 = 99\%$). The changes in mAlb were statistically different between the two groups ($N = 270$, MD $= -42.53$, 95% CI: $-65.48$–$-19.58, Z = 3.63, P = 0.0003$). We speculated that the source of heterogeneity might have been the different test method of mAlb, such as radioimmunoassay or enzyme-linked immunosorbent assay.

### Blood Lipid

This meta-analysis evaluated two blood lipid indicators, LDL-C and TG. Four studies reported that the LDL-C indicator had low heterogeneity ($\chi^2 = 3.02, P = 0.39, I^2 = 1\%$), and a fixed-effect model was adopted. It was determined that the LDL-C level of treatment groups was lower than that of the control groups (MD $= -0.39$, 95% CI: $-0.58$–$-0.20, Z = 4.05, P < 0.0001$) (Figure 9).

Nine studies reported the TG indicator and lacked heterogeneity ($\chi^2 = 51.42, P < 0.00001, I^2 = 84\%$). Therefore, a fixed-effects model was used for data analysis. The results revealed that the TG level in treatment groups was lower than that in control groups ($N = 811$, MD $= -0.39$, 95% CI: $-0.63$–$-0.15, Z = 3.21, P = 0.001$) (Figure 10). The differences in TG between FFXST plus western medicine and control groups, subgroup analysis was performed. Two of the studies compared changes in TG in FFXST monotherapy treatment with that of control groups, and the results revealed there was no statistical difference TG improvement between the two groups ($N = 198$, MD $= -0.36$, 95% CI: $-1.15$–$-0.44, Z = 0.87, P = 0.38$) with heterogeneity ($\chi^2 = 7.78, P = 0.005, I^2 = 87\%$). The difference between the two studies may have been caused by the difference in intervention measures in the control groups. The control group in one study (Peng S. L., 2015) used valsartan 80 mg QD (daily), whereas the control group in the other study (Wang M. L., 2017) used a blank control.

### Glycemic Control

In this review, we mainly assessed HbA1c. This indicator was reported in eight studies, involving 671 people. A fixed-effect model was used because there was no heterogeneity between studies ($\chi^2 = 4.5, P = 0.72, I^2 = 0\%$). The results indicated that there was no significant difference in HbA1c between two groups in the meta-analysis (MD $= -0.02$, 95% CI: $-0.12$–$-0.08, Z = 0.31, P = 0.76$) (Figure 9).
Figure 3: Risk of bias summary.
### Table 1: Study Comparison and Meta-Analysis Results

<table>
<thead>
<tr>
<th>Study or subgroup</th>
<th>Experimental</th>
<th>Control</th>
<th>Weight (%)</th>
<th>Mean difference</th>
<th>Mean difference</th>
</tr>
</thead>
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<tr>
<td></td>
<td>Mean SD</td>
<td>Total</td>
<td>Mean SD</td>
<td>Total</td>
<td>IV, random, 95% CI</td>
</tr>
<tr>
<td>1.11.1 FFXST alone</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Wang ML 2017</td>
<td>112.6 26.5</td>
<td>40</td>
<td>164.2 29.4</td>
<td>38</td>
<td>22.0</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>-51.60 [-64.04, -39.16]</td>
</tr>
<tr>
<td>Heterogeneity: not applicable</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Test for overall effect: Z = 8.13 (P &lt; 0.00001)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1.11.2 FFXST plus ACEI/ARB</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Dai XM 2012</td>
<td>59.24 46.32</td>
<td>40</td>
<td>85.46 49.32</td>
<td>40</td>
<td>18.6</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>-26.22 [-47.19, -5.25]</td>
</tr>
<tr>
<td>Wang XN 2012</td>
<td>57.6 21.8</td>
<td>38</td>
<td>93.5 32.2</td>
<td>42</td>
<td>22.2</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>-35.90 [-47.85, -23.95]</td>
</tr>
<tr>
<td>Yun P 2013</td>
<td>124.8 17.2</td>
<td>48</td>
<td>136.4 15.3</td>
<td>49</td>
<td>23.8</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>-11.60 [-18.08, -5.12]</td>
</tr>
<tr>
<td>Heterogeneity: tau² = 173.41; ch² = 13.44, df = 3 (P = 0.004); I² = 78%</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Test for overall effect: Z = 3.09 (P = 0.002)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Subtotal (95% CI)</td>
<td>150</td>
<td>155</td>
<td>78.0</td>
<td>-24.61 [-40.21, -9.02]</td>
<td></td>
</tr>
<tr>
<td>Heterogeneity: tau² = 356.35; ch² = 36.88, df = 4 (P &lt; 0.0001); I² = 89%</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Test for overall effect: Z = 3.31 (P = 0.0009)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total (95% Cl)</td>
<td>190</td>
<td>193</td>
<td>100.0%</td>
<td>-30.98 [-49.30, -12.66]</td>
<td></td>
</tr>
</tbody>
</table>

### Figure 5: Forest plots of FFXST effects on UAER.

### Figure 6: Forest plots of FFXST effects on ACR.

### Figure 7: Forest plots of FFXST effects on mAlb.
The difference in TG between the FFXST plus ACEI/ARB and control groups was compared in the other seven studies. The results indicated there was no statistical difference in TG improvement between the two groups ($N = 613, MD = -0.36$, 95% CI: $-0.59$ to $-0.13, Z = 3.11, P = 0.0002$) with heterogeneity ($\chi^2 = 24.17$, $P = 0.0005$, $\text{I}^2 = 75\%$). Sensitivity analysis showed that heterogeneity decreased after the removal of two studies (Dai XM, 2012, and Yun P, 2013) ($\chi^2 = 3.12$, $P = 0.08$, $\text{I}^2 = 68\%$). The improvement in CRP was significantly different between the two groups ($N = 148, MD = -1.92$, 95% CI: $-2.64$ to $-1.20, Z = 5.22, P < 0.00001$) (Figure 13).

3.4.9. Inflammatory Index. Three studies assessed the inflammatory indexes, including CRP, HS-CRP, IL-6, and TNF-α. This review mainly evaluated CRP. Two studies reported the CRP indicator, with heterogeneity ($\chi^2 = 3.12$, $P = 0.08$, $\text{I}^2 = 68\%$). The improvement in CRP was significantly different between the two groups ($N = 148, MD = -1.92$, 95% CI: $-2.64$ to $-1.20, Z = 5.22, P < 0.00001$) (Figure 13).

3.4.10. Adverse Events. Three of the 13 RCTs mentioned adverse events in treatment groups. No adverse reactions occurred in the treatment group one study (Lu YZ, 2009).
3.4.11. Publication Bias. We could not conduct the funnel plot analysis for the detection of publication bias because of an insufficient number of experiments.

3.5. Grade Assessment. According to the GRADE, the quality of evidence was rated as moderate for the primary outcome, and the secondary outcomes were rated as low, very low, or moderate (Table 2).

4. Discussion

DKD is one of the common microvascular complications of DM. Early prevention and treatment can delay the occurrence and progression of DKD, which is of great significance.
Table 2: GRADE assessment of quality of evidence for outcomes.

<table>
<thead>
<tr>
<th>Quality assessment</th>
<th>Summary of findings</th>
<th>Risk of bias</th>
<th>Inconsistency</th>
<th>Indirectness</th>
<th>Imprecision</th>
<th>Overall quality of evidence</th>
<th>Study event rates (%)</th>
<th>Relative effect (95% CI)</th>
<th>Anticipated absolute effects</th>
</tr>
</thead>
<tbody>
<tr>
<td>Participants (studies) follow-up</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total effective rate (CRITICAL OUTCOME)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>●●●● MODERATE due to risk of bias</td>
<td>56/89 (62.9%)</td>
<td>76/88 (86.4%)</td>
<td>RR 1.37 (1.15 to 1.64)</td>
</tr>
<tr>
<td>UAER (IMPORTANT OUTCOME; better indicated by lower values)</td>
<td>629 per 1000</td>
<td>Serious¹</td>
<td>No serious inconsistency</td>
<td>No serious indirectness</td>
<td>No serious imprecision</td>
<td>Undetected</td>
<td>56/89 (62.9%)</td>
<td>76/88 (86.4%)</td>
<td>233 more per 1000 (from 94 more to 403 more)</td>
</tr>
<tr>
<td>ACR (IMPORTANT OUTCOME; better indicated by lower values)</td>
<td>627 per 1000</td>
<td>Serious¹</td>
<td>Series²</td>
<td>No serious indirectness</td>
<td>No serious imprecision</td>
<td>Undetected</td>
<td>193</td>
<td>190</td>
<td>The mean acr in the intervention groups was 30.98 lower (49.3 to 12.66 lower)</td>
</tr>
<tr>
<td>mAlb (mg/L) (IMPORTANT OUTCOME; better indicated by lower values)</td>
<td>548 (6 studies)</td>
<td>Serious¹</td>
<td>Series²</td>
<td>No serious indirectness</td>
<td>No serious imprecision</td>
<td>Undetected</td>
<td>318</td>
<td>320</td>
<td>The mean mlab (mg/l) in the intervention groups was 36.29 lower (54.45 to 18.13 lower)</td>
</tr>
<tr>
<td>BUN (mmol/L) (IMPORTANT OUTCOME; better indicated by lower values)</td>
<td>338 (4 studies)</td>
<td>Serious¹</td>
<td>Series²</td>
<td>No serious indirectness</td>
<td>Serious³</td>
<td>Undetected</td>
<td>167</td>
<td>171</td>
<td>The mean bun (mmol/l) in the intervention groups was 0.59 lower (1.46 lower to 0.27 higher)</td>
</tr>
<tr>
<td>HbA1C (IMPORTANT OUTCOME; better indicated by lower values)</td>
<td>671 (8 studies)</td>
<td>Serious¹</td>
<td>No serious inconsistency</td>
<td>No serious indirectness</td>
<td>Serious³</td>
<td>Undetected</td>
<td>337</td>
<td>334</td>
<td>The mean hba1c in the intervention groups was 0.02 lower (0.12 lower to 0.08 higher)</td>
</tr>
<tr>
<td>LDL-C (IMPORTANT OUTCOME; better indicated by lower values)</td>
<td>353 (4 studies)</td>
<td>Serious¹</td>
<td>No serious inconsistency</td>
<td>No serious indirectness</td>
<td>No serious imprecision</td>
<td>Undetected</td>
<td>175</td>
<td>178</td>
<td>The mean ldl-c in the intervention groups was 0.39 lower (0.58 to 0.2 lower)</td>
</tr>
<tr>
<td>TG (IMPORTANT OUTCOME; better indicated by lower values)</td>
<td>811 (9 studies)</td>
<td>Serious¹</td>
<td>Series²</td>
<td>No serious indirectness</td>
<td>No serious imprecision</td>
<td>Undetected</td>
<td>407</td>
<td>404</td>
<td>The mean tg in the intervention groups was 0.39 lower (0.63 to 0.15 lower)</td>
</tr>
<tr>
<td>SBP (NOT IMPORTANT OUTCOME; better indicated by lower values)</td>
<td>275 (3 studies)</td>
<td>Serious</td>
<td>Serious²</td>
<td>No serious indirectness</td>
<td>Serious³</td>
<td>Undetected</td>
<td>138</td>
<td>137</td>
<td>The mean shp in the intervention groups was 1.26 lower (3.86 lower to 1.34 higher)</td>
</tr>
<tr>
<td>CRP (mg/L) (NOT IMPORTANT OUTCOME; better indicated by lower values)</td>
<td>148 (2 studies)</td>
<td>Serious¹</td>
<td>Series²</td>
<td>No serious indirectness</td>
<td>No serious imprecision</td>
<td>Undetected</td>
<td>74</td>
<td>74</td>
<td>The mean crp (mg/l) in the intervention groups was 1.92 lower (2.64 to 1.2 lower)</td>
</tr>
</tbody>
</table>

(1) The blind method of the included study was not mentioned. (2) Significant heterogeneity, I² = 88%. (3) Significant heterogeneity, I² = 91%. (4) Significant heterogeneity, I² = 99%. (5) 95% CI crosses the invalid line. (6) Significant heterogeneity, I² = 84%. (7) Significant heterogeneity, I² = 58%. (8) Significant heterogeneity, I² = 68%.
to the improvement, survival rate, and quality of life of diabetic patients [29]. Clinical practice shows that TCM has the characteristics of multitarget, multipathway, and low adverse reaction [30–32] and has a great potential in the intervention of DKD. FFXST is a Chinese patent medicine approved by the State Food and Drug Administration of China and has been widely used for the treatment of DKD. This meta-analysis suggests that FFXST may be a safe and effective treatment for DKD. This meta-analysis revealed that FFXST for DKD was superior to the treatments provided the control group in total effective rate. Additionally, FFXST exhibited advantages in improving proteinuria indicators (UAER, ACR, and mAlb), blood lipid (LDL-C, TG), and inflammatory index (CRP), but not in lowering BUN or HbA1c levels or blood pressure in DKD patients. However, it has been reported that FFXST has a certain antihypertensive effect, which needs to be confirmed by further studies in the future [33].

There were many limitations to our meta-analysis. Although the dosage and form of FFXST used in the 13 studies were consistent, the treatment periods, DM type, and DKD stage of patients were not similar among the RCTs. Furthermore, the methodological quality of the studies was generally low, and the sample size was not reported in the 13 RCTs. All of the previously mentioned factors might have negatively affected the reliability of the research results.

Regarding the outcome, only two in the 13 RCTs reported the total effective rate, which was the primary outcome in our review. The improvement of the symptoms of patients is an important part of the evaluation of the efficacy of DKD treatment, but only one study (Lu YZ, 2009) reported the change of symptom score of patients before and after treatment. Thus, we could not evaluate the improvement effect of FFXST on the symptoms of patients. It was necessary to standardize the DKD efficacy evaluation system in clinical trials, which could have improved the reliability of the analysis. One study (Dai XM, 2012) also showed that FFXST could improve hemorheology. However, such reports were rare, and more pharmacological and clinical studies are needed to verify the mechanism of FFXST in the treatment of DKD. Clinical events are often recommended as primary outcome indicators for clinical studies; however, no trials assessed the incidence of DKD clinical endpoint events (death/entry to ESRD) or other adverse indicators in our study, which may not be conducive to explain the effect of FFXST for DKD. The GRADE results showed that the evidence quality of the total effective rate and LDL-C level was moderate, and the quality of the remaining outcomes was low or very low. This is mainly due to the fact that most of the included studies did not use blind methods or the large heterogeneity between studies. Therefore, more rigorous clinical studies are still needed to confirm the efficacy of FFXST in the treatment of DKD.

Follow-up and adverse event reports were insufficient among the 13 studies, with only three studies reporting adverse events and one study reporting a follow-up; thus, this meta-analysis was unable to assess the long-term efficacy and safety of FFXST for DKD.

5. Conclusions

Our meta-analysis suggested that the Chinese patent medicine FFXST was superior to that of the treatment of the control group in the improvement of total effective rate, reduction of proteinuria, and lowered blood lipid. DKD patients, especially who are in the stage of Mogensen III, accompanied by abnormalities in indicators of UAER, ACR, mAlb, LDL-C, TG, and CRP, can be treated with FFXST or combined with western medicine. However, FFXST may not be an optimizing option to improve abnormal indicators of SBP, HbA1c, and BUN in DKD patients.

However, the long-term efficacy and safety of FFXST for DKD is uncertain because most studies included in this review were of low quality, having small sample sizes and high heterogeneity. Thus, high-quality, large-scale, and multicenter RCTs are needed to validate the current results.

Data Availability

All data generated or analyzed during the study are available and included in this published article.

Disclosure

YMT and JH are the co-first authors.

Conflicts of Interest

All authors declare that there are no conflicts of interest.

Authors’ Contributions

YMT, JH, and QN conceived the meta-analysis; QW, YZ, WDS, YTZ, and HG performed data collection and analysis. YMT and JH wrote and revised the manuscript. All authors approved the final version of the paper. YMT and JH contributed equally to this work.

Acknowledgments

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References

Evidence-Based Complementary and Alternative Medicine


