Xiaokeping mixture inhibits diabetic nephropathy in streptozotocin-induced rats through blocking TGF-β1/Smad7 signaling

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Background: Diabetic nephropathy (DN) is a major cause of chronic kidney failure and characterized by excessive deposition of extracellular matrix. Evidence have shown that transforming growth factor-β1 (TGF-β1) is a key mediator in the development of DN. However, treatment of DN by blocking the TGF-β1/Smad7 pathway remains limited. Xiaokeping mixture (XKP), a traditional Chinese herbal compound, has been used for treatment in patients with DN for many years.

Methods: In the present study, TGF-β1/Smad7 pathway analysis was used to evaluate the therapeutic effect of XKP on DN rats induced by streptozotocin and to address the underlying molecular mechanism. Male rats were divided into four groups: normal control, untreated control group (fed with high fat), irbesartan-treated DN, and XKP-treated DN, respectively. Levels of serum creatinine, blood urea nitrogen, urine protein of 24 hours, and triacylglycerol were detected. Pathological changes of renal tissues were observed by hematoxylin–eosin staining. Immunohistochemical and Western blot analysis were used to detect the expressions of TGF-β1 and Smad7.

Results: The results demonstrated that XKP can effectively reduce the levels of glucose, serum creatinine, blood urea nitrogen, urine protein of 24 hours, and triacylglycerol. Further studies indicated that inhibition of DN in XKP-treated DN rats was associated with inhibition of TGF-β1/Smad7 signaling as demonstrated by downregulation of TGF-β1 but upregulation of Smad7.

Conclusion: The data obtained from the present study indicate that XKP may be a therapeutic agent for DN.

Keywords: Xiaokeping mixture, diabetic nephropathy, transforming growth factor-beta, Smad7

Introduction

Diabetic nephropathy (DN) is a major cause of chronic kidney failure and characterized by excessive deposition of extracellular matrix (ECM). Persistent ECM production can be accelerated with severe renal injury, which results in large amounts of fibrinous tissue are generated and thus a vicious circle is formed progressively. So, it is essential to identify appropriate pharmacologic interventions to prevent renal tubulointerstitial fibrosis, especially to improve recovery of ECM following renal injury. Transforming growth factor-β1 (TGF-β1) signaling is a well-recognized pathway leading to the development of DN. A typical role of TGF-β1 is its biologic effects can exert through the Smad protein signaling pathways. Thus, inhibiting the TGF-β1/Smad signaling pathway is helpful for preventing renal tubulointerstitial fibrosis and preserving renal function.
For thousands of years, traditional Chinese medicines (TCMs) have played an important role in health maintenance for peoples throughout the world. Xiaokeping mixture (XKP) is a TCM preparation developed from a famous TCM doctor, Mr Kuijun Shi (approved by the Food and Drug Administration of Zhejiang province, medical license H20100002). It composed of Astragalus membranaceus, Rhizoma dioscoreae, Radix rehmanniae preparata, Radix ophiopogonis, radices trichosanthis, and chrysanthemum. XKP has been used for the treatment of diabetic mellitus for many years. Our previous studies have shown that XKP could decrease fasting blood glucose levels, increase insulin sensitivity index, etc. However, as TCMs usually operate in vivo through multi-components, multi-ways, and multi-targets, the molecular mechanisms of XKP remain unclear. In the present study, we sought to determine whether XKP has therapeutic potential for DN and investigated underlying mechanisms of its action in rats with accelerated diabetic kidney.

Methods

Animal and experimental protocols

All experimental procedures were conducted in accordance with the ethics committee of Tongde Hospital of Zhejiang province, and in compliance with the National Institutes of Health Guide for the Care and Use of Laboratory Animals. Male Sprague–Dawley rats with weights ranging from 180 to 220 g were purchased from the Shanghai SLAC Laboratory Animal CO. LTD. The rats were housed in an air conditioned room at 24°C–25°C, humidity of 65%–69% under a 12-hour dark/light cycle, and were given food and water freely. After 1 week adaptation, the rats were divided into a normal control group (NC, n=12) that was fed a standard diet and a high-fat (HF) group that received HF diet (containing 67.5% standard laboratory rat chow, 15% lard, 15% sugar, 2% cholesterol, and 0.5% bile salts). After 4 weeks, rats on the HF diet were treated with a single intravenous injection of 60 mg/kg streptozotocin (Sigma-Aldrich, Inc., St Louis, MO, USA). Animals were considered to be diabetic if they had plasma glucose concentrations of 16.7 mmol/L or greater, in addition to polyuria and other diabetic features. All rats were randomly divided into three groups as follows (n=12 each group): (1) untreated control group (fed with HF, HF group); (2) irbesartan-treated group (an ARB as known positive control, irbesartan-treated DN [IRB-DN]; 17.5 mg/kg/day, diluted in 0.5% carboxymethyl cellulose); and (3) XKP-treated DN (XKP-DN, 1.2 g/kg/day, diluted in 0.5% carboxymethyl cellulose). All drugs were administered via intra-gastric gavage once a day for 16 weeks.

Biochemical analysis and light microscopy

Body weight of rats was measured after treatments for 16 weeks, and blood of rats was sampled from the tail vein, the blood glucose levels were measured by a One Touch blood glucose monitoring system. And metabolic cages were used to collect urine samples of 24 hours, the urinary protein excretion was determined by the Bradford method. The total cholesterol, triglycerides, urea nitrogen, and serum creatinine were detected by an automatic biochemistry analyzer. Tissue for light microscopy was fixed in 4% (w/v) paraformaldehyde and then embedded in paraffin. Four micrometer thick sections of renal tissue were processed for hematoxylin–eosin and Masson’s trichrome staining.

Western blot analysis

Renal proteins were fractionated by sodium dodecyl sulfate polyacrylamide gelelectrophoresis (SDS-PAGE) and transferred onto nitrocellulose difluoride membranes. The membranes were blocked in 5% bovine serum albumin for 1 hour and incubated with a primary antibody at 4°C overnight, then they were washed in Tris-buffered saline, 0.1% Tween 20, and incubated with secondary antibody at 20°C for 1 hour. After washing, signals were visualized with an enhanced chemiluminescent system. Protein levels were normalized against β-actin.

Statistical analysis

The data were presented as the mean± standard deviation. Statistical analysis was carried out using SPSS 19.0. The differences among multiple groups were evaluated by one-way analysis of variances. P<0.05 was considered as statistically significant.

Results

Animal characteristics

At the beginning of study, all of the rats behaved normally. During the experiment, the rats in the HF diet group gradually showed listlessness, polyuria, and lost weight. After 7 weeks, the rats in the HF group appeared symptoms such as ulcer and cataracts. After 16 weeks of drug therapy, the symptoms mentioned above improved significantly both in IRB-DN group and in XKP-DN group.

Effects of XKP on bodyweight, blood glucose, and 24 hours urinary protein

At the end of study, two rats died in the HF group and all rats were alive in the other groups. After 16 weeks of drug therapy, rats in NC group displayed progressive gain of body weight whereas rats in HF group showed less gain of body weight. However, either in XKP-DN group or in IRB-DN group the
drug did not prevent the reduction in body weight, as shown in Table 1. Rats with untreated DN showed severe hyperglycemia characterized by elevated fasting and postprandial plasma glucose levels \(P<0.01\). However, treatment with XKP decreased plasma glucose levels significantly, when compared with those of HF group rats \(P<0.05\) (Table 2). The blood urea nitrogen, serum creatinine, and urinary total protein increased significantly in comparison with those of NC group rats \(P<0.01\). However, treatment with both XKP and irbesartan reduced the ratio of kidney to body weight and reversed renal dysfunction and proteinuria \(P<0.05\).

As shown in Table 3, rats with untreated DN showed severe dyslipidemia. The serum total cholesterol, triglycerides, and low-density lipoprotein cholesterol; levels have increased significantly, when compared with those of NC group rats \(P<0.01\). However, treatment with XKP and irbesartan alleviated hyperlipidemia significantly in rats with DN \(P<0.05\).

### Effects of XKP on renal structure

As shown in Figure 1, glomerular and tubular structures were examined by periodic acid–Schiff staining, there were no obvious abnormalities in glomerular and tubular structures of the kidneys in rats of NC group. However, there was a significant mesangial expansion, thickened basement membranes, and occluded capillaries in rats of HF group, and there was a significant collagen deposition and fibrosis in the tubulo-interstitium of rats. However, both XKP and irbesartan significantly reduced the changes in both glomerular and tubular structures.

### Effects of XKP on TGF-β1 and Smad7 expression

As shown in Figure 2, there were positive particles of TGF-β1 protein expression in rats of NC group (Figure 2A), with a little of tan grain. Whereas there were a lot of tan particles in rats of HF group (Figure 2B, compare with Figure 2A, \(P<0.01\)). Compare with HF group, there were much little particles in IRB-DN group (Figure 2C, \(P<0.05\)) or in XKP-DN group (Figure 2D, \(P<0.05\)), suggesting that TGF-β1 protein expression have increased on the DN rat kidney tissue; however, both XKP and irbesartan significantly reduced the TGF-β1 protein expression on the DN rat kidney tissue.

In the cytoplasm, there were positive of Smad7 protein expression in rats of NC group (Figure 2E), with a lot of tan particles, and the color is deep. Whereas there were much little tan particles in rats of HF group (Figure 2F, compare with Figure 2E, \(P<0.01\)). Compare with HF group, there were much more tan particles in IRB-DN group (Figure 2G, \(P<0.05\)) or in XKP-DN group (Figure 2H, \(P<0.05\)), suggesting that TGF-β1 protein expression have increased on the DN rat kidney tissue; however, both XKP and irbesartan significantly increased the Smad7 protein expression on the DN rat kidney tissue.

We next investigated the mechanisms by which XKP treatment attenuated DN by studying the TGF-β1/Smad7 signaling pathway. TGF-β1/Smad7 signaling was highly activated in the diabetic kidney as revealed by a marked upregulation of TGF-β1, but lower levels of an inhibitory Smad7 protein expression level, as shown in Figure 3. Treatment with XKP significantly reduced the TGF-β1 protein expression on the DN rat kidney tissue, which was associated with significantly increased Smad7 protein expression.

### Table 1 Effect of XKP on weight/kidney index in rats with DN at 16th week

<table>
<thead>
<tr>
<th>Group</th>
<th>Body weight (g)</th>
<th>Kidney/body weight (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>NC</td>
<td>492±39.7</td>
<td>0.757±0.02</td>
</tr>
<tr>
<td>HF</td>
<td>204±47.7</td>
<td>1.89±0.36</td>
</tr>
<tr>
<td>IRB-DN</td>
<td>284±28.7</td>
<td>1.35±0.32</td>
</tr>
<tr>
<td>XKP-DN</td>
<td>285±42.9</td>
<td>1.29±0.35</td>
</tr>
</tbody>
</table>

**Notes:** *P<0.01* compared with NC rats and *P<0.05* compared with HF rats. Data shown as mean ± standard deviation.

**Abbreviations:** DN, diabetic nephropathy; NC, normal control; XKP, Xiaokeping mixture; XKP-DN, XKP-treated DN

### Table 2 Effect of XKP on GLU and renal function in rats with DN at 16th week

<table>
<thead>
<tr>
<th>Group</th>
<th>GLU (mmol/L)</th>
<th>BUN (mmol/L)</th>
<th>SCR (μmol/L)</th>
<th>Urinary total protein (mg/24 h)</th>
</tr>
</thead>
<tbody>
<tr>
<td>NC</td>
<td>5.20±3.69</td>
<td>5.57±0.82</td>
<td>56.59±1.22</td>
<td>18.25±2.28</td>
</tr>
<tr>
<td>HF</td>
<td>24.33±5.04</td>
<td>15.02±1.32</td>
<td>92.51±1.15</td>
<td>92.24±2.46</td>
</tr>
<tr>
<td>IRB-DN</td>
<td>11.44±3.35</td>
<td>8.24±1.79</td>
<td>78.63±1.94</td>
<td>52.82±2.81</td>
</tr>
<tr>
<td>XKP-DN</td>
<td>9.82±4.24</td>
<td>8.32±0.81</td>
<td>80.44±1.85</td>
<td>55.09±3.07</td>
</tr>
</tbody>
</table>

**Notes:** *P<0.01* compared with NC rats and *P<0.05* compared with HF rats. Data shown as mean ± standard deviation.

**Abbreviations:** BUN, blood urea nitrogen; NC, diabetic nephropathy; DI, glucose; HF, high fat; IRB-DN, irbesartan-treated DN; NC, normal control; SCR, serum creatinine; XKP, Xiaokeping mixture; XKP-DN, XKP-treated DN; h, hours.

### Table 3 Effect of XKP on plasma lipid profiles in rats with DN at 16th week

<table>
<thead>
<tr>
<th>Group</th>
<th>TG (mmol/L)</th>
<th>TC (mmol/L)</th>
<th>LDL-C (mmol/L)</th>
<th>HDL-C (mmol/L)</th>
</tr>
</thead>
<tbody>
<tr>
<td>NC</td>
<td>1.14±0.22</td>
<td>2.07±0.28</td>
<td>0.59±0.15</td>
<td>1.55±0.28</td>
</tr>
<tr>
<td>HF</td>
<td>2.82±1.55</td>
<td>7.92±1.42</td>
<td>4.64±1.55</td>
<td>1.24±1.22</td>
</tr>
<tr>
<td>IRB-DN</td>
<td>1.73±0.42</td>
<td>5.24±1.92</td>
<td>1.73±0.27</td>
<td>1.47±0.32</td>
</tr>
<tr>
<td>XKP-DN</td>
<td>1.74±0.46</td>
<td>4.92±0.84</td>
<td>1.74±0.28</td>
<td>1.49±0.35</td>
</tr>
</tbody>
</table>

**Notes:** *P<0.01* compared with NC rats and *P<0.05* compared with HF rats. Data shown as mean ± standard deviation.

**Abbreviations:** DN, diabetic nephropathy; NC, normal control; XKP, Xiaokeping mixture; XKP-DN, XKP-treated DN; Tc, total cholesterol; TC, total cholesterol; TG, triglycerides.
Figure 1. Effects of XKP on renal pathology of experimental rats. 
Notes: Representative pictures of PAS staining (original magnification ×400) are presented as (A–D). (A) NC group; (B) HF group; (C) IRB-DN; and (D) XKP-DN. 
Abbreviations: DN, diabetic nephropathy; HF, high fat; IRB-DN, irbesartan-treated DN; NC, normal control; PAS, periodic acid–Schiff; XKP, Xiaokeeping mixture; XKP-DN, XKP-treated DN.

Figure 2. Effects of XKP on TgF-β1 and Smad7 expression of experimental rats. 
Notes: Representative pictures of PAS staining (original magnification ×400) are presented as (A–H). (A and E) NC group; (B and F) HF group; (C and G) IRB-DN; and (D and H) XKP-DN. 
Abbreviations: DN, diabetic nephropathy; HF, high fat; IRB-DN, irbesartan-treated DN; NC, normal control; PAS, periodic acid–Schiff; TgF-β1, transforming growth factor-β1; XKP, Xiaokeeping mixture; XKP-DN, XKP-treated DN.
Discussion

TCM formulas are an important source for the development of new drugs to prevent and treat diabetes and its complications. In the present study, we examined the therapeutic effect and the underlying mechanisms of XKP in a rat model of DN. We found that administration of XKP significantly inhibited streptozotocin-induced kidney fibrosis and markedly decreased proteinuria. The inhibitory effect of XKP on diabetic kidney disease was associated with inactivation of TGF-β1/Smad7 signaling.

DN is characterized by pathophysiological changes in glomerular hyperfiltration, renal hypertrophy, tubular function and then progress to proteinuria and reduction in glomerular filtration rate. Many studies have demonstrated that TGF-β1 plays an important role in experimental diabetic kidney disease and human DN. TGF-β1 is a key profibrotic cytokine that acts by stimulating synthesis of ECM components, decreasing matrix degradation, and promoting cell–matrix interactions. Moreover, TGF-β1 is also involved in tubule glomerular sclerosis and podocyte apoptosis in diabetes. Smad7, an inhibitory Smad, can block TGF-β1/Smad signaling by binding to Smurf2 to form an E3 ubiquitin ligase that targets the TGF-β1 receptor as well as Smads including Smad7 for degradation. Once Smad7 is degraded, activation of Smad3 and renal fibrosis is enhanced. The functional importance of TGF-β1/Smad signaling in DN is demonstrated in a number of animal models, in which overexpression of Smad7 inhibits diabetic renal injury. In our present study, treatment with XKP attenuated diabetic renal injury by rebalancing the TGF-β1/Smad signaling pathway. A similar mechanism was also evident by a positive treatment control with irbesartan. Thus, inhibition of TGF-β1/Smad7 signaling could be an important mechanism, by which XKP attenuated diabetic kidney disease.

Practitioners of TCM believe that in the early stage of DN, patients mostly exhibit deficiency in conjunction with dryness-yang in lung and spleen. This was manifested by frequent urination and easily getting thirsty, soreness and weakness of waist and knees, dizziness, and tinnitus. During the course of DN, the yin deficiency spreads to the yang, causing a yin-and-yang deficiency. XKP has long been used for the treatment of “kidney-Yang” deficiency syndrome. Accumulating evidence from the ethnomedical aspect indicates that the herbs and their chemical constituents are beneficial for the recovery of renal function, for example, A. membranaceus, also known as Huangqi in People’s Republic of China, has been frequently used in TCM for several centuries. As one of the 50 fundamental herbs, it is traditionally employed for the treatment of diabetes, for wound healing and strengthening immunity. It also possesses anti-inflammatory, hepatoprotective, antioxidative, and cardioprotective properties.

Conclusion

In the present study, we performed an analysis to evaluate the therapeutic effect of XKP treatment on DN rats and address the underlying molecular mechanism. Our results indicate that XKP treatment could ameliorate DN symptoms and inhibiting glucose and lipid metabolism. The mechanism is closely related to inhibit TGF-β1/Smad7 signaling, which further underscoring the potential clinical benefits of XKP in the treatment of DN.

Acknowledgments

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Disclosure

The authors report no conflicts of interest in this work.

References