ABSTRACT: To investigate the effect of total anthraquinone of Polygonum cuspidatum (TAPC) in diabetic rats which were established by streptozotocin. Method: The diabetic rat model was established by injected with streptozotocin, and then randomized into 5 groups. Which were fed with high dosage TAPC group, medium dosage TAPC group, low dosage TAPC group (400mg•kg$^{-1}$,200mg•kg$^{-1}$,100mg•kg$^{-1}$), metformin group (208 mg•kg$^{-1}$) and model group. Control group was set up with healthy normal rats. All the rats were administrated once a day for 30 days. Blood glucose (BG) in the limosis were measured on the 20, 30 day and then rats were sacrificed. The levels of insulin, insulin antibody, leptin, C-peptide in orbital blood were measured. Results: Diabetic model rats successfully. Compared with model group, The high dosage TAPC group can significantly reduce the level of blood suga on 20 day($P<0.05$), Can significantly reduce the level of blood suga on 30 day($P<0.01$), Each dosage TAPC group can significantly reduce the level of C-peptide ($P<0.01$). Conclusion: TAPC have good hypoglycemic effect on the diabetic rats established by injection of streptozotocin KEYWORD: Total anthraquinone of Polygonum cuspidatum (TAPC); Diabete model; Streptozotocin; Insulin antibody; Leptin; C-peptide.

1 INTRODUCTION

Polygonum cuspidatum, also known as "Kowloon root", "Yin Yang Lin", tepid, bitter, liver, gall bladder, lung, as Polygonum cuspidatum Sieb.et Zucc of dried roots and rhizomes, which as the medicine first appeared in "Thunder Gun Sunburn"[1]. The main effect is expelling wind and dampness, stasis and pain, cough and phlegm. Modern research shows that Polygonum cuspidatum contains anthraquinone compounds, flavonoids, water-soluble polysaccharides, tannins and trace elements and other components[2]. The total anthraquinone of Polygonum cuspidatum, including emodin ether, stilbene compound, which also contains tannin and polysaccharide[3].Modern pharmacological study have found that aqueous extract of Polygonum cuspidatum $\alpha$-glucosidase was significantly inhibited[4]; Polygonum cuspidatum had been reported that anthraquinone compounds for hypoglycemic active ingredients[5]; Another report the tannins of Polygonum cuspidatum has good hypoglycemic effect on the alloxan diabetic mice[6]. Polygonum cuspidatum as promoting blood circulation to remove blood stasis hypoglycemic drugs in folk had application [7]. In order to investigate the characteristics of Polygonum cuspidatum and effect on the treatment of diabetes, the experimental to investigate the effect of total anthraquinone of Polygonum cuspidatum on streptozotocin-induced diabetic rats model.

2 MATERIALS

2.1 Animals

Wistar rats, male, whose weight were 180-200g, were supplied by the Experimental Animal Center of Hebei Province (Animal permit number: 701022).

2.2 The experimental reagents and drugs

The TAPC, provided by the Henan University of Traditional Chinese Medicine chemistry department, content more than 50%; Metformin Hydrochloride Tablets was from Shanghai Pharmaceutical Group Co.,Ltd. Xinyi Pharmaceutical Factory; Streptozotocin (STZ) was from sigma company; Physiological saline, zhengzhou yonghe pharmaceutical co., LTD; Citric acid (AR) was from
Hubei Pharmaceutical company Bose station; Glucose kit was from Zhejiang East Ou Biological Engineering Co., Ltd. Insulin radiation immunoassay kit, c-peptide radiation immunoassay kit and insulin antibody radiation immunoassay kit were all from Beijing Kemei Doya Biological Technology Co., Ltd.

2.3 The experimental instrument
UV-2000 UV-visible spectrophotometer, UNICO (Shanghai) Instrument Co., Ltd; Constant temperature water bath, Beijing Guangming Instrument Factory; FA (N) /JA (N) series electronic balance, Shanghai Minqiao Precision Instrument Co., Ltd; Adjustable liquid shifter, Shanghai Leibo Analysis Instrument Co., Ltd.

3 METHODS
100 Wistar male rats. After fasting 12h, Random take 90 rat formulated at pH 4.2 ) after fasting 12h and the remaining as the blank group, Tail vein injection of an equal volume of citrate buffer[8]. On the tenth day, tail blood test blood sugars, select 50 rats which BG>11.1 mmol·L⁻¹, and obviously drinking, eating, urinating, then were fed with high, medium and low dosage TAPC suspension (400mg·kg⁻¹, 200mg·kg⁻¹, 100mg·kg⁻¹, 20mg·ml⁻¹, 10mg·ml⁻¹, 5mg·ml⁻¹, 2ml·100g⁻¹), metformin suspension (208mg·kg⁻¹, 20.8mg·ml⁻¹, 2ml·100g⁻¹), model group and blank control group were given the same volume physiological saline. Once a day, treated for 30 days.

4 STATISTICAL ANALYSIS
Data analysis used SPSS 13.0 for windows for statistical treatment. The differences of measurement data between groups were analyzed using ANOVA, ranked data used Ridit test.

5 RESULTS
5.1 Impact on blood sugar in the STZ rat model
From table 1, compared with the blank group, the level of BG in tenth, twentieth, thirtieth day was significantly increased (P<0.01), shows that the model successfully. Compared with the model group, in twentieth, thirtieth day, the medium dosage TAPC group and metformin group could remarkably degrade the level of BG (P<0.01); In thirtieth day, the high dosage TAPC group could remarkably degrade the level of BG (P<0.01); In twentieth day, the high, dosage TAPC group could obviously degrade the level of BG (P<0.05); Low dosage TAPC group has a tendency to lower the level of BG.

<table>
<thead>
<tr>
<th>Group</th>
<th>Dose (mg·kg⁻¹)</th>
<th>The level of BG (mmol·L⁻¹)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Began to glucose</td>
<td>10d</td>
</tr>
<tr>
<td>Blank group</td>
<td>4.82±0.85**</td>
<td>4.98±0.766**</td>
</tr>
<tr>
<td>Model group</td>
<td>15.47±2.600</td>
<td>17.55±2.492</td>
</tr>
<tr>
<td>Metformin group</td>
<td>15.87±0.936</td>
<td>15.66±2.090</td>
</tr>
<tr>
<td>High dosage TAPC group</td>
<td>15.66±2.805</td>
<td>16.90±3.266</td>
</tr>
<tr>
<td>Medium dosage TAPC group</td>
<td>16.23±3.019</td>
<td>16.47±1.860</td>
</tr>
<tr>
<td>Low dosage TAPC group</td>
<td>15.91±2.811</td>
<td>17.62±3.238</td>
</tr>
</tbody>
</table>

Table 1. Effect of TAPC on blood sugar in the STZ rat model ( X ± s, n = 10 )

Note: Compared with the model group, *P<0.05, **P<0.01

5.2 Impact on Serum insulin, C-peptide and leptin levels in the STZ rat model
From table 2, Compared with the control group, the model group could significantly degrade levels of insulin and C-peptide levels (P<0.01), significantly increase leptin levels (P<0.01). It suggests that the model copied successfully. Compared with the model group, medium dosage TAPC group and metformin group could significantly increase serum insulin levels (P<0.01), high dosage TAPC group could significantly increase serum insulin levels (P<0.01); Medium and low dosage TAPC group can significantly increase serum C-peptide levels (P<0.01); High, medium and low dosage TAPC group can significantly degrade serum leptin levels (P<0.01).
Table 2. Effect of TAPC on Serum insulin, C-peptide and leptin levels in the STZ rat model (X ± s, n = 10)

<table>
<thead>
<tr>
<th>Group</th>
<th>Dose (mg·kg⁻¹)</th>
<th>Serum insulin (×10-3μIU·ml⁻¹)</th>
<th>C-peptide (ng·ml⁻¹)</th>
<th>Leptin (ng·ml⁻¹)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Blank group</td>
<td></td>
<td>47.280±8.471**</td>
<td>0.109±0.034**</td>
<td>0.645±0.180**</td>
</tr>
<tr>
<td>Model group</td>
<td></td>
<td>26.850±4.809</td>
<td>0.043±0.016</td>
<td>1.517±0.346</td>
</tr>
<tr>
<td>Metformin group</td>
<td>208</td>
<td>37.320±6.064**</td>
<td>0.061±0.017</td>
<td>0.849±0.469**</td>
</tr>
<tr>
<td>High dosage TAPC group</td>
<td>400</td>
<td>33.129±5.393*</td>
<td>0.054±0.016</td>
<td>0.970±0.430**</td>
</tr>
<tr>
<td>Medium dosage TAPC group</td>
<td>200</td>
<td>36.887±5.077**</td>
<td>0.101±0.021**</td>
<td>0.630±0.235**</td>
</tr>
<tr>
<td>Low dosage TAPC group</td>
<td>100</td>
<td>29.927±5.884</td>
<td>0.074±0.017**</td>
<td>0.840±0.273**</td>
</tr>
</tbody>
</table>

Note: Compared with the model group, *P<0.05, **P<0.01

6 DISCUSSION

Diabetes in traditional Chinese medicine belong to the category of "excessive", and obviously drinking, eating, urining, thin body, or turbid urine, urine has a sweet taste; Modern medicine that diabetes is a chronic metabolic disease caused by a variety of causes, its basic pathology for the absolute or relative lack of insulin secretion and peripheral tissue is not sensitive to insulin, or cause is given priority to with sugar metabolic disorders, including fat and protein metabolism disorder of a systemic disease[9]. The experimental model by intravenous injection of STZ diabetic model making[10], because of its damaging effects on islet β cell in experimental animals is highly selective, and its mechanism induced diabetic animal model is first islet β cell in specific sites of DNA base alkylation further action ribosome synthesis in ADP enzyme and thus directly damage pancreatic β cell by NO and O2- - radicals two paths, making the islet β cell necrosis, insufficient insulin secretion, which can be successfully established diabetes model that is closer to human diabetes, has been widely used. Insulin, serum, C-peptide is an important indicator of insulin secretory function of cells; Leptin is a hormone gene from obesity produced, and insulin resistance are closely related [11]. Serum insulin, C-peptide is an important index reflecting cell secretion of insulin. Leptin is obese gene (OB) control coding, a cycle of endocrine hormone secreted by fat cells, which is one of the symbols of the direct effect of obesity and its main functions of the central nervous system leptin receptors, inhibiting the synthesis of neuropeptide Y, then to reduce appetite, reduce the intake of energy, and the relationship between leptin and insulin resistance is closely; As there is no cross reaction c-peptide and insulin antibodies, are not affected by the interference of insulin antibodies, its value can more directly, more objectively and more accurately reflects the function of islet beta cells [12].

This study shows that the TAPC can significantly degrade blood glucose in the STZ rat model, there was no significant difference between its hypoglycemic effect of metformin; and in hypoglycemic while significantly elevated levels of serum insulin and C-peptide levels significantly lower leptin levels which elevated the role of C-peptide levels significantly better than metformin group; but the regulation of insulin antibodies was not obvious. The result suggest that the TAPC in Polygonum cuspidatum has good curative effect on streptozotocin induced diabetic rat model, can stimulate the secretion of islet C-peptide and insulin, improve the symptoms of diabetes. This experiment is used in the treatment of diabetes Polygonum cuspidatum provided data support and support, but also for the further development and utilization of Polygonum cuspidatum foundation.

7 ACKNOWLEDGEMENTS

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