Protective role of berberine and *Coptischinensis* extract on T2MD rats and associated islet Rin-5f cells

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Abstract. The aim of the present study was to compare the different effects of berberine (Ber) and *Coptischinensis* extract (CCE) on a rat model of type 2 diabetes mellitus (T2DM), and the islet Rin-5f cell line was used to examine the differences between Ber and CCE and the underlying mechanisms. CCE was extracted and purified prior to analysis. Male Sprague-Dawley rats were provided with a high-fat diet to induce insulin resistance prior to injecting with streptozotocin to establish the T2DM model. The T2DM rats were treated with Ber and CCE, and blood samples and pancreatic tissues were obtained and compared to examine T2DM metabolic syndromes among the groups of rats, which included healthy rats, model rats, and model rats treated with Ber and CCE at different doses between 0 and 8 weeks. The protective effects of Ber and CCE on the Rin-5f islet cell line were also evaluated. The effects on Rin-5f cell proliferation and cell cycle, glucose-stimulated insulin release test (GSIS), the anti-apoptotic effects caused by fat induction, and protein expression levels of poly ADP-ribose polymerase (PARP-1) were evaluated. The results showed that the content of the prepared CCE was 96.07% for five alkaloids. When it was used for treatment of the T2DM rats, compared with Ber, metformin and rosiglitazone, the fasting blood glucose, glucosylated serum protein (GSP) and glucose infusion rate indices in the fasting rats were ameliorated, compared with those in the T2MD rats, with no significant differences between treatment with Ber or CCE and metformin or rosiglitazone. The indices of mean optical density and fasting β-cell function index (FBCI) were different following treatment with Ber and CCE, compared with those in the model rats, which may have stimulated the pancreatic secretion of insulin. When Ber and CCE were used to examine the protective effects on Rin-5f cells, it was found that the Rin-5f cell GSIS, cell cycle, lipotoxic islet cell proliferation and protein expression of PARP-1 were altered and improved, which may have protected pancreatic islet β-cells by improving islet β-cell proliferation and the protein expression of PARP-1. CCE and Ber exerted similar effects when used for the treatment of T2DM rats, and may have stimulated the pancreatic secretion of insulin through the protective effect on islet β-cells via improving islet β-cell proliferation and the protein expression of PARP-1.

Introduction

Berberine (Ber) is an isoquinoline alkaloid isolated from *Coptischinensis* Franch, which differs from *Coptischinensis* extract (CCE) (1). Studies have shown that Ber offers potential for the treatment of type 2 diabetes mellitus (T2DM) (2-7), whereas CCE is primarily one of the components of a Chinese traditional formula for the treatment of diabetes mellitus (DM) via the prevention of associated vascular and cardiac damage, renal fibrosis and lipid metabolism (8-10). Ber is a classical natural medicine, the applications of which have attracted interest in investigations focused on the treatment of DM (11-15). The pharmacological effects of Ber on glucose metabolism...
have been shown to involve pathways, including improving insulin resistance (16), promoting insulin secretion (17-19), protecting diabetic nephropathy (20-22), modulating gut microbiota (23,24), affecting phospholipid metabolites (25), preventing cardiac dysfunction in T2DM (26-28), inhibiting gluconeogenesis in the liver (29,30), reducing glucose absorption and enhancing glucose consumption (31,32), ameliorating hepatosteatosis and stimulating glycolysis (33,34), and regulating lipid metabolism (35,36). The potency of Ber in the treatment of DM is predominantly associated with its antioxidant and anti-inflammatory activities (37,38), and few studies have focused on the pancreatic-derived factor of islet β-cells (39-41). The protein expression of poly ADP-ribose polymerase (PARP)-1 in islet β-cells may also be an important pathway for DM treatment (42).

Therefore, the present study selected Ber and CCE as the subjects for investigation, and their different effects on T2DM rats were examined. The indices of fasting blood glucose (FBG), glucosylated serum protein (GSP) and glucose infusion rate (GIR) in fasting rats were compared among treatment groups of rats, and the pancreatic indices of mean optical density (MOD) and fasting β-cell functional index (FBCI) were compared for the evaluation of insulin secretion. Ber and CCE were also used to investigate their protective effects on RIN-5f cells, in which results of the glucose-stimulated insulin release test (GSIS), cell cycle, lipotoxic islet cell proliferation and protein expression levels of PARP-1 were analyzed. CCE and Ber possessed equal effects in the treatment of T2MD rats, which may have stimulated the pancreatic secretion of insulin. In the RIN-5f cells, the GSIS, protection of lipotoxic islet cell proliferation and apoptosis, and the protein expression levels of PARP-1 were improved, which may protect pancreatic of islet β-cells by improving islet β-cell proliferation and the protein expression of PARP-1.

Materials and methods

Preparation and analysis of CCE. The herbal medicine Coptischinensis Franch (Guangzhou Baiyunshan Zhong Yi Pharmaceutical Co., Ltd., Guangzhou, China; batch no. 20090208) was extracted by the addition of 0.5% sulfuric acid (1:8 by weight) three times at a temperature of 100°C (1.5 h each time). The filtrates were mixed and concentrated by evaporation at 75°C to obtain a condensed solution (relative density of 1.07 at 55°C), following which the solution was filtered and the precipitate discarded prior to adjusting the pH to 9.0 with calcium hydroxide (10% by weight). The solution was then filtered and the pH was adjusted to 1.5 with hydrochloric acid (AR), following which sodium chloride (AR) was added to adjust the concentration (10% by weight). The solution was agitated to mix, prior to standing for 24 h, following which the precipitate was obtained and washed with water until a pH of 5.0 was obtained. The precipitate was refined three times using hydrochloric acid, sodium chloride (10% by weight) and water, according to the above-mentioned procedure. The obtained precipitate was evaporated to dryness at 60°C to obtain 8 kg of CCE.

The sample analysis was performed on an Agilent 1200 high-performance liquid chromatography (HPLC) system (Agilent Technologies, Inc., Santa Clara, CA, USA) equipped with a diode array detector at a wavelength of 345 nm. The HPLC column used was a Venusil XBP-C18 (Agela Technologies, Inc., Wilmington, DE, USA) analytical column (250x4.6 mm, 5 µm). The mobile phase composition was acetonitrile and 0.05 M potassium dihydrogen phosphate buffer solution (25:72, v/v) at a flow rate of 1.0 mL/min. The test solution was prepared by dissolving CCE in hydrochloric acid-methanol (1:100, v/v). The standard was prepared by dissolving the standard sepiberberine (EPI), coptisine (COP), palmatine (PAL), jatrorrhizine (JAT) and Ber in hydrochloric acid-methanol (1:100, v/v) to obtain appropriate concentrations for CCE sample analysis. The EPI, COP, PAL, JAT and Ber standards were purchased from the National Institutes for Food and Drug Control (Beijing, China) and Shanghai Yuanye Biotechnology Co., Ltd. (Shanghai, China).

Animals, model establishment and treatment. A total of 180 male Sprague-Dawley rats (90±10 g; 4 weeks old) of SPF grade were purchased from Vital River Lab Animal Technology Co., Ltd (Beijing, China), as was the high-fat diet, which contained 18% lard, 20% sucrose, 3% egg yolk and 59% basal diet (24.0% protein, 3.5% lipids and 60.5% carbohydrate). All experiments were approved by the Ethical Committee of Guang'anmen Hospital, China Academy of Chinese Medical Sciences (Beijing, China). The animals were cared for in accordance with the Guide for the Care and Use of Laboratory Animals of the National Institutes of Health (Bethesda, MD, USA). The rats were randomly divided into two groups; in the normal control group (n=20), rats were fed with a basal diet; in the high-fat diet group (n=160), rats were fed with the high-fat diet for a total of 8 weeks. The rat body weights were compared between the control group and the experimental group in eight randomly selected rats, and no significant differences in body weights were found between the two groups. The rats were randomly selected for eight hyperinsulinemiceuglycemic clamp experiments to assess insulin resistance, the results of glucose homeostasis and glucose infusion rate (GIR) are shown in Table I, with insulin resistance observed in the high-fat diet group rats. When the rats in the high-fat diet group showed significant insulin resistance, the rats were administered with an intravenous injection of streptozotocin (dissolved in 0.1 M citrate buffer in an ice bath at a concentration of 1%; pH 4.4) at a dose of 30 mg/kg to establish the T2DM rat model. The rats in the normal diet group were administered with an intravenous injection of an equal volume of citric acid-sodium buffer solution. After 1 week, the blood glucose levels of the rats were randomly measured, and a blood glucose level of >16.7 mM was considered to indicate successful model establishment. The model rats were randomly divided into eight groups according to blood glucose level and body weight. The rats in the control group and model group were orally administered with distilled water every day; rats in the rosiglitazone group received rosiglitazone at a dose of 0.4 mg/kg/d; rats in the metformin, Ber and CCE groups were administrated with metformin, Ber and CCE at a dose of 100 mg/kg/d (Table II), respectively. The rats in each group were administered with distilled water or drugs for 8 weeks.

Measurement of fasting blood glucose and glycosylated serum protein. The rats were fasted overnight for 12 h (8.00 p.m-8.00
a.m.) with free access to water. The FBG was measured the following morning using a blood glucose meter via the tail vein. The blood serum was separated by centrifugation at 4,000 x g for 15 min at 4°C to determine the levels of glycated serum protein (GSP).

**Hyperinsulinemic-euglycemic clamp experiment.** The rats were fasted overnight prior to the hyperinsulinemic-euglycemic clamp experiment with free access to water. The rats were anesthetized via 2% sodium pentobarbital intraperitoneal injection, the dose of which was calculated by body weight at 40 mg/kg. The right carotid artery and the left jugular vein were isolated and incubated with a silica gel epidural catheter (inner diameter, 0.6 mm; outer diameter, 1 mm), which contained saline heparin (50 U/ml) to maintain patency. Arterial blood glucose was determined using a blood glucose meter. The venous catheter was connected to the tee and the vein connected to the outlet, with the inlet end connected to infusions of insulin (infusion rate 8 mU/kg/min, insulin albumin diluted with 0.5% bovine serum from Gibco, Thermo Fisher Scientific, Inc. (Waltham, MA, USA; lot: 1527494) and glucose (20%) using a microcomputer digital micro-injection syringe pump from the jugular vein. The blood glucose levels of the rats were determined following of 30 min of insulin and glucose infusion. The blood was obtained from the jugular vein to measure glucose values. Firstly, short-acting porcine insulin (Sigma-Aldrich; Merck Millipore (Darmstadt, Germany; 1:500; batch no. 20110925) was constantly infused, and blood glucose was determined once every 10 min. When the blood glucose level exceeded the baseline value ± 0.5 mM, 20% glucose was infused to adjust the blood GIR on the basis of the control value ± 0.5 mM. The GIR was adjusted in the shortest possible time in accordance with a normal blood glucose level. The procedure was repeated until three consecutive blood glucose values were stable within the above range; the steady state value of the three average GIR values (mg·kg⁻¹·min⁻¹) was obtained.

**Pancreatic tissue immunohistochemistry.** The rats were sacrificed under anesthesia. The pancreatic tissue was isolated and embedded in paraffin prior to being cut into sections (2x2 cm, 8 μm) for insulin immunohistochemistry (INS). The tissue sections were stained with guinea pig anti-porcine insulin antibody (cat. no. 129-10332; RayBiotech, Inc., Norcross, GA 30092; 1:100) in the β-cells. For the second step of the DAKO antibody Envision™ system (Glostrup, Denmark), DAB was used as a color reagent for the β-cell tissue sections. The Envision procedure was as follows: Tissue sections were skinned prior to hydration by rinsing in distilled water and were placed in TBS, to which with 0.3% hydrogen peroxide was added to block endogenous peroxidase, and then incubated for 15 min at 4°C. The tissue sections were rinsed in distilled water and placed into TBS for rinsing for 5 min, which was repeated three times. Following incubation for 2 h at 4°C, TBS rinsing was performed for 5 min three times, Envision™ was added prior to incubation for 60 min, followed by rinsing in TBS for 5 min three times. The DAB source chromogenic solution was used for staining for 3 min, and color development was terminated with distilled water. The sections were then dehydrated in alcohol. A negative control experiment was also performed, in which TBS was used instead of primary antibody as a negative blank control. The results were observed under a light microscope (Nikon Corporation, Tokyo, Japan). The MOD was obtained from Sigma Photo Pro software version 5.5.1 (Sigma-Aldrich; Merck Millipore) by counts. The FBCI was calculated based on fasting insulin level (FINS) and FBG using the following formula: FBCI = FINS / FBG.

**Cell culture.** The Rin-5f cells were cultured in medium comprising fetal bovine serum (FBS) diluted in 1640 culture medium (1:10, v/v). When ~80% of the Rin-5f cells had adhered to the bottom of the dish, the original culture medium was discarded and replaced with 5% BSA in 1640, and cultured for 36 h for synchronized cultivation. The Rin-5f cell line was provided by the Cell Library of Wuxi Medical School, Jiangnan University (Wuxi, China). The 1640 culture medium, FBS and antibiotic-antimycotic were purchased from Invitrogen; Thermo Fisher Scientific, Inc. The kit used for western blot analysis was purchased from GE Healthcare Life Sciences (Chalfont, UK). PARP-1 antibody (cat. no. AE90773Mu) was purchased from Cell Signaling Technology, Inc. (Danvers, MA, USA); HRP-labeled goat anti-mouse IgG (H+L) (cat. no. SN133) were all obtained from NanJing KeyGen Biotech Co., Ltd. (Nanjing, China), β-actin (cat. no. P0068) were obtained from Beyotime Institute of Biotechnology (Shanghai, China). The FITC Annexin V Apoptosis Detection Kit II was purchased from BD Biosciences (Franklin Lakes, NJ, USA). Fatty acid-free bovine serum albumin (FFA) and other chemical reagents were purchased from Sigma-Aldrich; Merck Millipore. FFA was dissolved with 0.1 M NaOH solution at 70°C, and then shaken for 10 min prior to filtering through a 0.22-μm filter as the stock solution at a concentration of 100 mM. The FFA solution was dissolved in deionized water to obtain a stock solution concentration of 50 g/l at 55°C, prior to filtering through a 0.22-μm filter. A solution was then prepared, which contained 5% FBS and 0.25 mM FFA in 1640 culture medium. A Synergy 2 microplate reader (BioTek Instruments, Inc., Winooski, VT, USA) was used to measure the absorbance. The mRNA expression levels were detected using the line-general-time fluorescence quantitative polymerase chain reaction system (Line-Gene 9600; Hangzhou Bior Technology Co., Ltd., Hangzhou, China). An RNA prep pure cell kit (cat. no. DP430) was obtained from Tiangen Biotech Co., Ltd. (Beijing, China). PCR was performed using a Verso 1-step QRT-PCR kit Plus ROX vial (cat. no. AB-4100/A; Thermo Fisher Scientific, Inc.), which contained 50 μl Verso

<table>
<thead>
<tr>
<th>Group</th>
<th>Glucose homeostasis (mM/l)</th>
<th>Glucose infusion rate (mg·kg⁻¹·min⁻¹)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control</td>
<td>5.0±0.1</td>
<td>23.38±1.26</td>
</tr>
<tr>
<td>High-fat diet</td>
<td>5.0±0.2</td>
<td>20.27±1.60a</td>
</tr>
</tbody>
</table>

Data are presented as the mean ± standard deviation (n=8). *P<0.05, vs. control.
**Table II. Details of the rats in each experimental group.**

<table>
<thead>
<tr>
<th>Group</th>
<th>n</th>
<th>Blood glucose (mM)</th>
<th>Body weight (g)</th>
<th>Dose (mg/kg/d)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control</td>
<td>20</td>
<td>4.8±1.1*</td>
<td>369.6±19.6</td>
<td>Distilled water</td>
</tr>
<tr>
<td>Model</td>
<td>15</td>
<td>23.0±3.8</td>
<td>372.2±30.4</td>
<td>Distilled water</td>
</tr>
<tr>
<td>Metformin</td>
<td>15</td>
<td>23.1±3.7</td>
<td>370.8±39.1</td>
<td>100</td>
</tr>
<tr>
<td>Rosiglitazone</td>
<td>15</td>
<td>23.0±3.9</td>
<td>365.1±42.9</td>
<td>0.4</td>
</tr>
<tr>
<td>Ber-low</td>
<td>15</td>
<td>23.3±3.7</td>
<td>362.1±40.6</td>
<td>50</td>
</tr>
<tr>
<td>Ber-middle</td>
<td>15</td>
<td>23.1±3.9</td>
<td>362.9±39.3</td>
<td>100</td>
</tr>
<tr>
<td>Ber-high</td>
<td>15</td>
<td>23.2±4.1</td>
<td>363.1±45.5</td>
<td>250</td>
</tr>
<tr>
<td>CCE-low</td>
<td>15</td>
<td>23.1±3.8</td>
<td>355.8±54.7</td>
<td>50</td>
</tr>
<tr>
<td>CCE-middle</td>
<td>15</td>
<td>23.2±3.9</td>
<td>359.8±51.1</td>
<td>100</td>
</tr>
<tr>
<td>CCE-high</td>
<td>15</td>
<td>23.1±3.9</td>
<td>344.9±50.9</td>
<td>250</td>
</tr>
</tbody>
</table>

*P<0.05 (control group vs. CCE or Ber). Ber, berberine; CCE, Coptischinensis extract.

Enzyme mix, 50 µl yellow RT Enhancer, a 1-step QRT-PCR mix (2X 1.25 ml) and 25 µl brown ROX dye. The fluorophore used in the reaction was Taqman (cat. no. 4304437; Thermo Fisher Scientific, Inc.). The 1-step QRT-PCR mix contained Thermo-Start™ DNA polymerase (cat. no. AB-4100/A; Thermo Fisher Scientific, Inc.), which had 3' to 3' polymerization and exonuclease activity but lacked 3' to 5' exonuclease activity. The kit used for cDNA synthesis and reverse transcription were from the Verso 1-step QRT-PCR kit Plus ROX vial, where the Verso™ Enzyme mix (cat. no. AB-4100/A; Thermo Fisher Scientific, Inc.) contained the reverse transcriptase enzyme for long cDNA strand generation. and also contained RNase inhibitor to protect RNA templates from degradation (according to the manufacturer's protocol). Cycling conditions were as follows (according to the kit protocol): 50°C for 15 min for one cycle, thermostatic activation at 95°C for 15 min for one cycle, followed by 40 cycles at 95°C for 15 sec and extension at 60°C for 60 sec (43). Primers were as follows: forward, 5'GCC CTAAAGGCTCAAGACGAC3' and reverse, 5'CACCACTGGC ATCAGCTACTG3' for PARP-1; forward, 5'CACCATGCCTCTCACCTTTG3' and reverse, 5'CCACACCTGGTGC TGTAG3 for GAPDH. Image capture was performed using the Fluorochem 9800 system (Alpha Innotech). The 2^−ΔΔCq method was used for quantification (44).

**Rin-5f cell survival rate assay using MTT.** The Rin-5f cells were transferred into a 96-well plate at ~1x10^6 cells per well, following which different concentrations of the drugs were added to each well and incubated for 0, 24, 48 and 72 h prior to the MTT assay. Culture medium without Rin-5f cells and drug-free culture medium with Rin-5f cells were used as blank and control groups, respectively. Each group was examined in triplicate (n=3). The obtained absorbance values were used to calculate the Rin-5f cell survival rate (SR%). The absorbance (A) of each well was determined using a Synergy 2 micro-plate reader at wavelength of 570 nm, SR% was used to evaluate the effects of Ber and CCE on Rin-5f cells, calculated using the following formula: SR% = (A_{Drug groups} - A_{blank}) / (A_{Control group} - A_{blank}).

**GSIS release test.** The Rin-5f cells were induced with the drugs for 3 days, following which the drug was washed off, and 1.0 g/l glucose was added, prior to culture in 2% BSA 1640 culture medium in a CO₂ incubator at 37°C for 10 min. The supernatant was aspirated, and 3 g/l glucose was added to cells for culture in 2% BSA 1640 culture medium for 45 min. Again, the supernatant was discarded and the cells were washed twice with the culture medium. Subsequently, 1 g/l glucose was added and the cells were cultured in 2% BSA in 1640 medium for 45 min, following which the supernatant was discarded. The levels of insulin were measured using chemiluminescence, and the insulin release index was calculated by comparing the high glucose intake insulin value with the low glucose insulin value (n=6). The experimental groups composed of control group, glibenclamide group, Ber groups at doses of 5, 20, 100, 150 and 250 µM, and CCE group at doses of 5, 20, 100, 150 and 250 µM.

**Analysis of cell cycle.** The Rin-5f cells were transferred into a 6-well plate at ~1x10^5 in each well for culture until 80% of cells had adhered to the bottom of the plate. Culture synchronization was performed for 36 h prior to discarding of the original culture medium. The cells were divided into a control group (5% BSA in 1640 medium) and drug-induced groups (0.25 mM FFA, BerorCCE at a dose of 100 µM in 5% BSA in 1640 medium) with three wells for each group. Following drug intervention for 48 h at 37°C, the trypsin-digested cells were collected by centrifugation (1,350 x g) at 4°C for 15 min, and the suspended and adherent cells were collected and washed twice with pre-cooled PBS at 4°C. The cells were then resuspended with binding buffer to a cell concentration of 10^6/ml, and 100 µl of each sample cell suspension was collected and added to0 µl propidium iodide solution (PI) and 100 ng/ml RNA enzyme-binding buffer. The mixture was incubated at 37°C for 30 min in a water bath. Subsequently, 400 µl of the binding buffer was added into each sample fluid, and cell cycle and distribution analyses were performed on the samples using flow cytometry.

**MTT determination of the inhibition of lipotoxic islet cell proliferation.** The Rin-5f cells were seeded into a 96-well plate (20,000 per well), following cell cultivation for synchronization. Different concentrations of the drug and 0.25 mM palmitic acid, low glucose 1640 culture medium, and 2%
BSA, respectively, were added to the wells, with six parallel wells for each group. The MTT assay was performed using an MTT kit, according to the manufacturer’s protocol, following cultivation for 24 h. The experiments were repeated three times. The absorbance (A) values were used to calculated islet cell proliferation inhibition rate (PIR) as follows:

$$PIR = [1 - (A_d - A_b) / (A_c - A_b)] \times 100\%,$$

where $A_d$, $A_c$ and $A_b$ represented the drug induced group, control group and blank group, respectively. There were 13 groups, including the control group, Ber groups at doses of 2, 10, 50, 100, 250 and 500 µM and the CCE groups.

Protective effects of Ber and CCE on high fat-induced islet cell apoptosis. Coherst apoptotic Rin-5f cell staining was performed to examine apoptosis. Sterilized slides were placed into 6-well cell culture plates; with $5 \times 10^5$ islets cells in each well. Following incubation for the synchronization, when cells had adhered to the bottom of the plate, 1640 low glucose culture medium, 0.25 mM palmitate and 2% BSA were added to induce cell apoptosis, following which the drugs were added for 24 h prior to removal of the culture medium. Following washing with PBS three times, the slides were removed and stained according to the manufacturer's protocol of the coherst staining kit. The numbers of apoptotic cells were counted in 100 cells per field on each slide in five fields of view under a fluorescence microscope. The apoptotic rate was calculated as follows: Apoptotic rate = number of apoptotic cells / 500 x 100%. The experimental groups comprised Ber and CCE at doses of 2, 10, 50, 100, 250 and 500 µM respectively.

Determination of islet cell apoptosis. The high fat-induced Rin-5f cells were divided into a control group (culture medium), model group (0.25 mM FFA, 5% BSA, 1640), Ber (100 µM) and CCE (100 µM), prior to induction for 24 h. The cells (5x10^5 islets cells per well) were stained according to the Annexin V-FITC/PI double staining method. The culture medium was removed and the cells were washed twice with Dulbecco’s phosphate-buffered saline (D-PBS), following which 0.25% trypsin was added to digest the cells, and the cell suspension was collected by centrifugation at 4°C for 5 min at 1,350 x g to remove the medium. The precipitate was resuspended with 0.5 ml binding buffer, following which 5 µl Annexin V-FITC reagent and 5 µl PI were added prior to vortexing for 1 min. The mixture was incubated at room temperature for 20 min in the dark. The binding of Annexin V-FITC was measured using flow cytometry to obtain the V-positive cell number and the mean fluorescence intensity (excitation wavelength, 488 nm; emission wavelength, 530 nm). A total of 1.0 ml of sample was analyzed for positive expression rate detection and adjusted to 1x10^4 cells in each vial. The results were presented on a bivariate flow cytometry scatter plot. The lower left quadrant, the upper right quadrant and the lower right quadrant represent live cells (FITC-/PI-), necrotic cells (FITC+/PI+) and apoptotic cells (FITC+/PI-), respectively.

Western blot analysis of the protein expression of PARP-1 in apoptotic islet cells. The Rin-5f cells were seeded into a 100 mm petri-dishes at a concentration of 10^5 cells per well. When 80% of the cells had adhered to the bottom of the wells, culture synchronization was performed for 36 h, prior to discarding of the culture medium. The cells were divided
into a control group (5% BSA, 1640 medium), model group (0.25 mM FFA, 5% BSA, 1640 medium) and drug groups (Ber or CCE, 0.25 mM FFA, 5% BSA, 1640 medium), prepared at doses of 50, 100, 150 and 200 µM for Ber and for CCE. The cells were incubated for 24 h prior to digestion with protein lysate to extract proteins at 4°C. The protein concentrations were determined according to the Bradford method as follows: For each 50 µg protein sample, isolation was performed by adding 130 ml polyacrylamide and 1 ml SDS gel per liter, followed by transfer onto a cellulose acetate membrane, and then blocked at room temperature with tris-buffered saline with tween-20 (TBST) containing 50 g/l of skim milk for 1 h. Subsequently, specific antibody was added and incubated at 4°C overnight, following which the membrane was washed with TBST three times (10 min each). The membrane was then incubated with horseradish peroxidase-labeled goat anti-rabbit IgG (1:1,000, diluted in 0.01 M sodium phosphate and 0.25 M NaCl at pH 7.6) at room temperature for 1 h, and then washed with TBST three times (10 min each). A sensitizing

Table III. Hyperinsulinemic-euglycemic clamp test results of experimental groups.

<table>
<thead>
<tr>
<th>Group</th>
<th>n</th>
<th>Glucose (mM)</th>
<th>GIR (mg·kg⁻¹·min⁻¹)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control</td>
<td>6</td>
<td>5.0±0.1</td>
<td>24.01±2.26a</td>
</tr>
<tr>
<td>Model</td>
<td>6</td>
<td>4.9±0.2</td>
<td>9.45±1.47</td>
</tr>
<tr>
<td>Metformin</td>
<td>6</td>
<td>5.0±0.2</td>
<td>15.64±1.82b</td>
</tr>
<tr>
<td>Ber-high</td>
<td>6</td>
<td>5.1±0.3</td>
<td>16.65±2.48b</td>
</tr>
<tr>
<td>CCE-high</td>
<td>6</td>
<td>5.0±0.2</td>
<td>15.65±1.33b</td>
</tr>
</tbody>
</table>

*a P<0.01, vs. model. GIR, glucose infusion rate; Ber, berberine; CCE, Coptischinensis extract.*

Figure 3. Graph showing the changes in rat FBG in each group. Data are presented as the mean ± standard deviation. FBG, fasting blood glucose. *P<0.05, compared with the model group rats.

Figure 4. Graph showing the levels of GSP of rats in each group. Data are presented as the mean ± standard deviation. GSP, glucosylated serum protein. *P<0.05, compared with the model group rats.

Figure 5. Pancreatic tissue immunochemistry and MOD results of each group of rats. Data are presented as the mean ± standard deviation. *P<0.05 in the model group rats compared with the other groups. MOD, mean optical density.

Figure 6. Basal insulin levels in the rats in each group. **P<0.01, compared with the model group rats.

Figure 7. FBCI values of the rats in each group. *P<0.05, compared with the model group rats; **P<0.01, compared with the model group rats. FBCI, fasting β-cell function index.
A reagent kit was used to detect chemiluminescent substrate. The Flurochem 8900 image acquisition system and Alpha View software version 3.0 (Protein Simple Bioscience and Technology Co., Ltd., Shanghai, China) were used for image capture and scanning for densitometric analysis, respectively.

Statistical analysis. Statistical analysis was performed on SPSS 11.5 software (SPSS, Inc., Chicago, IL, USA). An independent-samples t-test was used for intergroup comparison. All data are presented as the mean ± standard deviation. P<0.05 was considered to indicate a statistically significant difference.

Results

CCE sample analysis. The CCE was analyzed according to the developed HPLC method, the contents of EPI, COP, JAT, PAL and Ber were 2.79, 3.11, 8.03, 8.02 and 74.12% respectively. The content of the five alkaloids was 96.07%. The contents,
chemical structures and HPLC chromatography results for CCE alkaloid determination are shown in Fig. 1.

**Body weights of the rats.** The body weights in the control group of normal rats gradually increased between 0 and 8 weeks, which was also observed in the metformin group rats. In the model group, the body weights initially decreased and then increased slowly, whereas the body weights gradually decreased in the CCE-high group and Ber-high group. In the rosiglitazone group, body weights initially increased and then decreased as in the CCE and Ber groups (Fig. 2). Where rat health was affected by severe T2DM, body weights decreased initially (0-4 weeks) between a stage of severe disease and recovery, and then increased gradually (5-8 weeks). In each drug group, no decreases in body weights were observed between 0 and 4 weeks in each, including the rosiglitazone group, due to the protective effects of the drug. The body weights of rats were increased in the metformin group. Metformin is a biguanide antihyperglycemic agent used for the treatment of non-insulin-dependent DM, which may explain the increase in body weight.

**FBG and GSP.** The FBG levels were measured in each group; the FBG in the control group of normal rats was normal during the 8 weeks at a range of 3.9-7.2 mM. The FBG levels in rats of the metformin and rosiglitazone groups were 17.4 and 19.1 mM at 8 weeks, which were significantly different, compared with the levels in the model group rats (P<0.05). The FBG levels in the rats of the CCE-high group and Ber-high group were 19.5 and 20.4 mM at 8 weeks, which were also significantly different, compared with levels in the model group rats (P<0.05). No differences in FBG were found in the CCE-high group and Ber-high group, compared with the metformin or rosiglitazone groups. The trend of FBG followed an initial increase (4 weeks) and subsequent decrease (8 weeks), with the exception of the control group. The decreases in FBG were associated with the increased doses of the CCE and Ber (Fig. 3), and the decreases in GSP were associated with the increased doses of CCE and Ber (Fig. 4).

**Hyperinsulinemic-euglycemic clamp test.** It was found that the GIR of the model group rats was significantly decreased, compared with the normal control group, metformin group, Ber-high group and CCE-high group rats in the hyperinsulinemic-euglycemic clamp experiment (P<0.01). Following drug intervention with metformin, Ber or CCE, GIR was significantly increased (P<0.01; Table III).

**Pancreatic tissue immunohistochemistry, insulin levels and FBGI.** The INS results showed that, in the control group of normal rats, pancreatic tissue was stained dark brown-yellow in color. The islet cells were expressed at a high level and were diffuse in the central region of the pancreatic tissue. In the model group rats, diffuse islet cells were observed in lower number, compared with the normal group, and were brown in color. The islet cells in the metformin, CCE-high group and Ber-high group rats were distributed in the central region of the pancreatic tissue and brown in color, the expression of which was high, compared with that in the model group rats. The MOD values obtained from each group were compared, and
the lowest MOD was recorded in the model group, which was significantly lower, compared with that of the normal control group (P<0.05). Following intervention with metformin, CCE or Ber, the MOD values were significantly higher, compared with that in the model group (P<0.05), however, no significant differences were found among the drug treatment groups (Fig. 5).

The basal insulin levels obtained from each group were also compared, the lowest of which was present in the model group rats (0.74±0.18), which was significantly (P<0.01) lower, compared with that in the control group rats (2.89±0.82), and was significantly higher, compared with those of the metformin, CCE or Ber drug intervention groups (P<0.01). No significant differences were found among three drug treatment groups (Fig. 6).

The FBCI values obtained from each group were also compared, the lowest of which was present in the model group rats (1.08±0.27), which was significantly (P<0.01) lower than that in the control group rats (6.14±1.09), and was significantly higher, compared with those in metformin, CCE or Ber drug intervention groups (P<0.05). No significant differences were found among the three drug treatment groups (Fig. 7).

 Rin-5f cell survival rate assay using MTT. According to the SR% formula, the SR% of Ber and CCE were determined at doses of 2, 10, 50, 100, 250 and 500 μM, respectively. The results are shown in Table IV. There was no effect on the survival rate of Rin-5f cells treated with Ber and CCE up to 500 μM.

 GSIS test on Rin-5f cells. The GSIS of Ber and CCE were determined at doses of 5, 20, 100, 150 and 250 μM (Table V). The results showed that Ber and CCE increased GSIS of the Rin-5f cells at a dose of 100 μM (P<0.05), compared with the control group. Ber and CCE promoted the secretion of insulin, however, this was to a lesser degree, compared with that of glibenclamide.

 Analysis of Rin-5f cell cycle. Following analysis of the cell cycle induced by 0.25 mM FFA prior to Ber and CCE, the G1 stage of the Rin-5f cell cycle was decreased, whereas the G2 and S cell cycle stages were increased (Fig. 8). No significant differences were found between the control group and Ber or CCE group.

 Effects of Ber and CCE on lipotoxic islet cell proliferation. An MTT assay was used to examine the effects of Ber and CCE on lipotoxic islet cell proliferation. The PIR was determined in each group (Table VI), which was decreased significantly by Ber and CCE at doses of 100 μM, compared with that in the control group (P<0.05). The effect of Beronlipotoxic islet cell proliferation was with dose-dependent at the dose range of 2-500 μM.

 Determination apoptosis of Rin-5f cells. Following induction of the Rin-5f cells with 0.10, 0.25 and 0.5 mM FFA for 24 h, the apoptosis of Rin-5f cells was examined at each period of time under a microscope. The cohest staining in the 0.5 mM FFA-induced group showed cell shrinkage into a round or oval shape, synapses had disappeared, cells were floating or suspended in medium with loss of adherent growth, and a large number of vacuoles were apparent with no visible nuclei. In the 0.25 mM FFA-induced group, there was partial cell shrinkage and adherent growth, and nuclei were visible. The numbers of apoptotic cells in each group were increased with increasing dose and time. The apoptosis of Rin-5f cells was determined using a Annexin V/FITC/PI double-labeled kit and flow cytometry to obtain the proportion of apoptotic Rin-5f cells in each group (Fig. 9). No significant difference in the proportions of apoptotic BRin-5f cells were observed in the groups at 6 h, however, apoptosis was significantly decreased following treatment with Ber or CCE at a dose of 100 μM for 24 h. There was a significant difference between the CCE group and control group (P<0.05).

Effects on islet cell protein expression of PARP-1. For each group, the grayscale of the protein expression ofPARP-1 was plotted for the Rin-5f cells at 24 h (Fig. 10), which revealed higher gray values in the sample groups, compared with the control group. The relative protein expression was determined and compared between the sample group and control group. The protein expression levels were significantly increased in the Ber and CCE groups at doses of 100, 200 and 500 μM. The dose of 200 μM had a more marked effect, compared with the other doses.

Discussion

The present study developed a simple preparation method to obtain CCE with a purity of 96.07% for alkaloids from Coptischinensis Franch. The systematic pharmacological function of the anti-DM treatments of Ber and CCE were investigated using a T2MD rat model and the Rin-5f cell line. When CCE was used to treat T2DM rats and compared with Ber at 8 weeks, the FBG of the rats was reduced. No significant differences were found between Ber or CCE and metformin or rosiglitazone; the FBG and GSP values were decreased significantly in all of the drug groups with dose-dependence. Ber and CCE increased GIR via improving insulin secretion.

When Ber and CCE were used to examine their effects on Rin-5f cells no toxicity towards the Rin-5f cells was observed; IRI and GSIS were increased, cell cycle was altered showing a decrease in G1 and an increase in G2+S. Ber and CCE exhibted a protective effect on lipotoxic islet cell proliferation, which was associated with the increased protein expression of PARP-1. CCE was more effective than Ber on altering the protein expression of PARP-1.

In conclusion, CCE and Ber possessed similar efficacies in the treatment of T2MD rats, the effects of which may have occurred through stimulating the pancreatic secretion of insulin, which may be associated with their protective role on islet β-cells by increasing islet β-cell proliferation and the protein expression of PARP-1.

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