Viability of primary cultured retinal neurons in a hyperglycemic condition

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Abstract
The retina of Wistar rats within 1–3 days of birth were dissociated into a retinal cell suspension using 0.05% trypsin digestion. The cell suspension was incubated in Dulbecco’s modified Eagle’s medium for 24 hours, followed by neurobasal medium for 5–7 days. Nissl staining showed that 79.86% of primary cultured retinal cells were positive and immunocytochemical staining showed that the purity of anti-neurofilament heavy chain antibody-positive cells was 71.53%, indicating that the primary culture system of rat retinal neurons was a reliable and stable cell system with neurons as the predominant cell type. The primary cultured retinal neurons were further treated with 0, 5.5, 15, 25, and 35 mM glucose for 24, 48, and 72 hours. The thiazolyl blue tetrazolium bromide test and flow cytometry showed that with increasing glucose concentration and treatment duration, the viability of retinal neurons was reduced, and apoptosis increased. In particular, 35 mM glucose exhibited the most significant effect at 72 hours. Thus, rat retinal neurons treated with 35 mM glucose for 72 hours can be used to simulate a neuronal model of diabetic retinopathy.

Key Words
neural regeneration; peripheral nerve injury; retina; neurons; apoptosis; hyperglycemia model; diabetic retinopathy; glucose; grants-supported paper; photographs-containing paper; neuroregeneration

Research Highlights
(1) Retinal neurons from neonatal Wistar rats (1–3 days) were cultured with plating medium and maintenance medium to establish a reliable and stable primary culture system using neurons as the predominant cells.
(2) The retinal neurons were treated with glucose at different concentrations for different time, and 35 mM for 72 hours was identified as the optimal concentration and intervention time.
(3) A retinal neuron model of high glucose was successfully established, which is significant for further studies of structure characteristics and biological functions of cell components in the retina, retinal pathological changes, and drug reactions.

INTRODUCTION
Retinal neurons are retinal photoreceptors and executors for nerve information transfer. Retinal functional damage caused by various causative factors is mainly manifested by loss of neuronal functions[1]. Thus, it is important to study physiologic and biochemical characteristics of retinal neurons in vitro. Isolation from the retina greatly damages neurons[2], so the isolation and culture of retinal neurons must be given careful attention. In addition, a large number
of studies have focused on cell-cell and cell-extracellular matrix interactions in the retina, but complex in vivo environmental effects greatly limits in vivo studies[5]. Therefore a stable and reliable in vitro culture system for retinal nerve cells is important for further understanding the structural characteristics and biological functions of components of various cells in the retina. In addition, it is important in the retina to understand the mechanisms of drug reactions and pathological conditions.

Diabetic retinopathy is a major oculopathy which can eventually lead to blindness. It is therefore a major focus and challenge of clinical and basic studies regarding oculopathy. Thus, it is clinically significant to study the pathogenesis and prevention of this disorder. Recent evidence has indicated that changes in retinal neurons and glial cells occur earlier than retinal capillary alterations in the initial stages of diabetes[4]. The changes in retinal neurons directly influence progression of diabetic retinopathy and may be the leading cause of diabetes-induced retinal capillary lesions[5]. Thus, understanding of retinal neuron changes is useful for prevention of diabetic retinopathy. In addition, in vivo experiments have been conducted in animal models to investigate the changes of retinal neurons in diabetic retinopathy[8]. Establishment of an in vitro retinal neuron model of high glucose to simulate the in vivo microenvironment of diabetes can eliminate extraneous interference of other factors in vivo, and allow direct observation of cell changes in specific stages of diabetes. This is an advantage over animal experiments.

Numerous studies have demonstrated that changes in retinal vascular endothelial cells[7-8], retinal pigment epithelium[9], retinal Müller cells[5], and retinal neurons[10-11] are highly correlated with the incidence and progression of diabetic retinopathy. A single in vitro cell model of high glucose injury could help elucidate the precise mechanism by which high glucose damages retinal cells. It would also provide experimental and theoretical evidence for prevention and treatment of diabetic retinopathy. There have been no reports of an in vitro cultured retinal neuron model of high glucose-induced injury. Takano et al[12] cultured retinal tissue blocks with 57 mM glucose for 3–10 days to investigate the influence of high glucose on axonal regeneration of retinal neurons from normal and streptozocin-induced diabetic mice, and found that high glucose suppressed axonal regeneration of retinal neurons in both groups. Another study established an apoptosis model using primary cultures of retinal neurons with 15 mM glucose for 9 days, to investigate the influence of erythropoietin, glucose, and nitric oxide on neurons[10]. Results showed that erythropoietin reduced nitric oxide levels and neuronal death in neuron cultures containing low concentrations of glucose. Santiago et al[11] utilized 25 mM glucose to culture retinal neurons in vitro for 7 days to establish a high glucose model, and found that neuronal apoptosis was associated with the caspase pathway. All the above retinal neuronal models of high glucose utilized different concentrations and durations of glucose treatment, providing a reference for our study. However, they were simply based on previously described methods, and did not continuously observe apoptotic retinal neurons in response to glucose treatment at different concentrations for different durations. Moreover, some of them did not utilize culture media specific for neurons. Thus, it is important for ophthalmology studies to develop an optimal method for in vitro culture of retinal neurons, to establish a stable and reliable model to study the effects of high glucose.

Our objective in this study was to establish an optimal culture system of retinal neurons, and a neuron model of high glucose, to establish a foundation for future studies of mechanisms by which drugs could protect retinal neurons in the presence of high glucose.

RESULTS

Morphology and phenotype of primary cultures of retinal neurons
The retina was harvested from Wistar rats within 1–3 days of birth for primary cultures of retinal neurons. After incubation for 30 minutes, the body of retinal cells was small and round shaped (Figure 1A). At 24 hours, the majority of cells adhered to the wall, and short processes emanated from some cells and accumulated at the center (Figure 1B). At 2–3 days, the processes were extended, about 1/2 to 1–2 folds longer than cell body length. The neurons were polygon- and oval-shaped with plump bodies, with surrounding visible nuclei and nucleoli (Figure 1C). After culturing for 5–6 days, the processes further enlarged and increased, accompanied by surrounding non-neuron cells (glial cells; Figure 1D). At 7–10 days, neurons continued growing, and the length of processes extended over 10-fold longer than cell bodies, gradually forming a complex network, with a gradual reduction in the number of non-neuron cells (Figure 1E). Up to 15 days, the processes of most cells became shortened, and neurons disintegrated or died (Figure 1F).
Nissl staining of retinal neurons cultured for 5–7 days showed a blue-violet stained cytoplasm, and granular Nissl bodies with clear structures. The cytoplasm of non-neuron cells was not stained, with light violet, round nuclei and clear nucleoli (Figure 2). The percentage of neurons was approximately 79.86%.

Immunocytochemical staining of retinal cells cultured for 5–7 days showed that 71.53% of cells were positive for anti-neurofilament antibody (Figure 3A), but only 21.13% were positive for anti-glial fibrillary acidic protein antibody (Figure 3B).
MTT assay showed that retinal neuron viability remained unchanged following 5.5 mM glucose treatment for 24–72 hours ($P > 0.05$). However, cell viability significantly decreased with increasing glucose concentration and intervention duration, compared with the control group (15 mM for 72 hours, $P < 0.05$; 25 mM for 48, 72 hours, $P < 0.05$; 35 mM for 48, 72 hours, $P < 0.01$). Most importantly, 35 mM glucose treatment for 72 hours significantly decreased cell viability ($P < 0.01$, Table 1).

Annexin/propidium iodide double staining was used to detect apoptosis. Flow cytometry showed that 5.5 mM glucose treatment for 24–72 hours did not influence apoptosis of retinal neurons ($P > 0.05$). However, the apoptosis rate was significantly increased with increasing glucose concentration and intervention duration compared with the control group (15 mM for 72 hours; 25 mM for 48, 72 hours; 35 mM for 48, 72 hours, $P < 0.05$ or $P < 0.01$). In particular, the apoptosis rate was significantly higher after 35 mM glucose treatment for 72 hours compared with the other groups ($P < 0.01$; Figures 4–6, Table 2).

### Discussion

Primary culture conditions of retinal neurons are controllable, allowing studies that cannot be conducted in vivo. Retinal neurons cannot divide, and can easily be contaminated by proliferating cells. Moreover, the conditions for retinal neuron survival are very specific. Thus, culture of retinal neurons is more difficult than other types of cells.[13]
Wistar rats within 1–3 days of birth, with an observable gap between neuroepithelium and pigment epithelium, allowing complete isolation of neuroepithelium from the pigment epithelium\textsuperscript{[14]}.

Thus, we used rats as experimental animals to obtain retinal neurons of high purity. (2) Time of sampling: generally, the survival rate of \textit{in vitro} cultured neurons is associated with the age of the donor animals\textsuperscript{[15]}. 

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**Figure 5**  Apoptosis of primary cultured retinal neurons exposed to glucose at different concentrations for 48 hours. (A) Normal control group; (B–E) 5.5, 15, 25, and 35 mM glucose groups. Flow cytometry detected and divided cells into four subgroups, including necrotic and non-viable apoptotic cells (annexin\textsuperscript{-}/propidium iodide (PI)\textsuperscript{+}) in the left upper quadrant, necrotic cells (annexin\textsuperscript{+}/PI\textsuperscript{+}) in the right upper quadrant, normal living cells (annexin\textsuperscript{-}/PI\textsuperscript{-}) in the left lower quadrant, and viable apoptotic cells (annexin\textsuperscript{+}/PI\textsuperscript{-}) in the right lower quadrant.

**Figure 6**  Apoptosis of primary cultured retinal neurons exposed to glucose at different concentrations for 72 hours. (A) Normal control group; (B–E) 5.5, 15, 25, and 35 mM glucose groups. Flow cytometry detected and divided cells into four subgroups, including necrotic and non-viable apoptotic cells (annexin\textsuperscript{-}/propidium iodide (PI)\textsuperscript{+}) in the left upper quadrant, necrotic cells (annexin\textsuperscript{+}/PI\textsuperscript{+}) in the right upper quadrant, normal living cells (annexin\textsuperscript{-}/PI\textsuperscript{-}) in the left lower quadrant, and viable apoptotic cells (annexin\textsuperscript{+}/PI\textsuperscript{-}) in the right lower quadrant.

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During isolation, neurons from embryos are in early development. The axons and dendrites have not extensively branched, nutritional dependence to target cells is low, and the epithelium of connective tissue is easily isolated[16]. However, it is difficult to confirm the age of the embryo and to obtain eyeball samples because of pigment epithelium contamination[17]. (3) Preparation of retinal cell suspension: the retinal cell suspension was prepared using a complete digestion method, so the quantity of obtained neurons depended on trypsin concentration and digestion duration. Following comparison of different methods, we utilized a method previously described by Santiago et al[11], which resulted in a large quantity of retinal neurons after retina tissues were digested with 0.05% trypsin at 37°C for 15 minutes. Low concentrations of trypsin or short durations of digestion resulted in nerve cell aggregation, while high concentrations of trypsin or long durations of digestion decreased cell viability. Our results demonstrated that enzyme digestion is a stable and reliable method for acquiring retinal neurons, providing a good experimental model for studying retinal disease. (4) Importance of adherence in cell growth and development: firm adherence is critical for neuron cultures, because the cell body can enlarge and the number of extended processes can increase. Takahashi et al[18] inoculated retinal ganglion cells on a glial cell monolayer, resulting in an increase in viability and an increase of neurite outgrowth. However, it is difficult to distinguish or isolate retinal ganglion cells and glial cells. Some studies also cultured cells on poly-L-Lysine- or poly-D-Lysine-pretreated glass slides to facilitate attachment[19-20]. In addition, a tectal extract of polylysine plus rat tail collagen increased adherence of cells compared with polylysine or rat tail collagen alone[21-22], and a tectal extract of cellular adhesion molecule L1, anti-Thy-1 monoclonal antibody, and vitronectin were also used[23]. The present study utilized polylysine (molecular weight 70 000)-coated glass slides. Polylysine can support neuronal growth and increase cell adherence and growth. In addition, Dulbecco’s modified Eagle’s medium containing newborn calf serum, F-12 nutrient mixtures, and glutamine were used, which provided sufficient nutrition and energy for neuronal adherence, to shorten the time of adherence to approximately 24 hours after inoculation. (5) Selection of maintenance medium: serum-containing Dulbecco’s modified Eagle’s medium has been frequently utilized for in vitro culture of neurons, generally supplemented with mitotic inhibitors (cytarabine) to prevent hyperplasia of glial cells and fibroblasts[24]. However, this medium is designed to supply the required nutrition for continuous proliferation, but is not suitable for neuronal growth. During isolation of cell suspensions of nerve tissues, glial cells, fibroblasts, and other non-neuron cells cannot be completely removed[25]. They are toxic to neurons, and could possibly lead to cell death[26]. Thus, the present study replaced Dulbecco’s modified Eagle’s medium with neurobasal medium/B27 serum-free supplements to maintain cultured cells that stably adhered. Serum-free neurobasal medium is utilized specifically for nerve cells. This medium can support purified neuronal growth, promote process differentiation and extension, and not allow continuous division or proliferation of glial cells and fibroblasts[27]. The medium also inhibits the effects of various components in the serum, resulting in neurons with active growth, high density and purity, but with a small number of non-neuron cells.

Table 2. Influence of glucose treatment at different concentrations on apoptosis rate (%) of retinal neurons

| Group       | Intervention time (hour) |
|-------------|--------------------------|
|             | 24          | 48          | 72          |
| Control     | 0.882±0.065 | 1.458±0.450 | 3.502±1.265 |
| 5.5 mM glucose | 0.925±0.055 | 3.538±1.301 | 4.095±1.948 |
| 15 mM glucose   | 4.497±2.221 | 10.280±4.380| 15.720±4.017a |
| 25 mM glucose    | 6.775±2.650 | 12.340±5.095a | 21.917±5.520a |
| 35 mM glucose    | 9.245±3.506 | 20.233±5.545b | 40.123±7.576b |
| F            | 16.561      | 21.606      | 62.335      |
| P            | 0.000       | 0.000       | 0.000       |

Results are expressed as mean ± SD of six parallel wells in each group. The experiments were performed in triplicate. *P < 0.05. **P < 0.01, vs. control group (one-way analysis of variance, intergroup multiple comparison using the Least Significant Difference test).

Apoptosis rate = [(number of annexin+propidium iodide (PI)+ cells) / (number of annexin+PI+ cells + number of annexin+/PI- cells)] × 100%.

Neurofilament is a cytoskeletal protein and a marker for mature axons. It is specifically distributed in the body and processes of neurons[28]. Neurofilament-positive staining can display the cell body, dendrites, and thick axons of neurons. Moreover, anti-neurofilament immunocytochemical staining has high sensitivity that can help visualize neuronal morphology[29]. In this study, the majority of retinal cells cultured for 5–7 days were positive for anti-neurofilament antibody, manifested as brownish-yellow particles in the cell body and processes, with darkly stained cell body and lightly stained processes. This indicated that neurofilament mainly accumulated in the cell body and processes, demonstrating that these cells have properties of retinal neurons[30], consistent with Nissl staining. Gliarial fibrillary acidic protein is specific to glial cells.
Anti-gliarial fibrillary acidic protein immunocytochemical studies showed that the percentage of gliarial fibrillary acidic protein antibody-positive cells was less than 30% of total retinal cells. This confirmed that the purity of the neurons was over 70%, supporting previous results\(^{31}\). Thus, this method can be used to culture retinal neurons with high purity, long survival time, and intact cell morphology, which provides conditions for studies of neuroprotection and drug action of retinal neurons.

High glucose is an important factor during onset of diabetic retinopathy. An earlier study showed that glucose at concentrations of 25–30 mM in an *in vitro* culture system is equivalent to a high glucose state of diabetes *in vivo*\(^{33}\). The MTT assay and flow cytometry showed that 5.5 mM glucose had minimal effect on neurons, but cell viability gradually decreased and apoptosis rate increased with prolonged glucose concentration and culture time. In particular, the effects of glucose treatment at 35 mM for 72 hours were the most significant. Preliminary studies showed that all cells almost died when cultured with glucose over 40 mM, consistent with the results of a previous study that retinal vascular endothelial cells were badly damaged in response to glucose over 80 mM\(^{35}\). This study utilized an annexin V/fluorescein isothiocyanate apoptosis detection kit, and flow cytometry demonstrated that apoptosis was the main injury to retinal neurons induced by high glucose. The early apoptosis rate was high in the 15–35 mM glucose treatment groups, possibly because either annexin V-propidium iodide detection was highly sensitive\(^{33}\) and/or reduced viability and increased apoptosis resulted from long durations of *in vitro* culture\(^{34}\). In addition, the cell source may influence the apoptosis rate. For example, a large number of neurons in neonatal rats apoptosed within one week after birth\(^{35-36}\). MTT and flow cytometry confirmed that 35 mM glucose treatment for 72 hours was the optimal concentration and intervention time for simulating the high glucose state *in vivo*.

Recent studies have utilized glucose at 15–33 mM to establish a neuron model of high glucose\(^{10-11}\). There are some differences in high glucose models due to the objectives of the studies, the culture media used, the purity of retinal neurons, and the varied experimental animals and conditions. For example, Dulbecco’s modified Eagle’s medium/F-12 culture media containing cytarabine to inhibit giall cell proliferation was used, but the glucose concentration was very low. In the present study, Neurobasal medium was used to culture retinal neurons for a short period of time (72 hours). As we did not use cytarabine, there were some glial cells, which could promote neuronal growth, partly attenuating the high glucose-induced injury. Thus, the glucose concentration used here was slightly higher than previous studies.

This also demonstrates interaction between neurons and glial cells under high glucose conditions. Additional studies may further focus on these relationships. Regarding the natural death of neurons, we established a normal control group and treated cells cultured for 3–4 days with glucose at different concentrations not more than 3 days. This can maintain cells in the best condition during experimentation, so that the experimental data were more relevant.

In summary, a retinal neuron model of high glucose was successfully established. This model can prevent overdose of glucose-induced severe neuronal injury, and inhibit natural death after long periods of intervention. Moreover, MTT and flow cytometry demonstrated neuron apoptosis in response to high glucose. It should therefore be feasible to study the neuroprotective role of drugs in early diabetic retinopathy. Furthermore, the results of this study provide novel ideas for treating neuronal changes in early diabetic retinopathy.

### MATERIALS AND METHODS

#### Design
Parallel study of cytology.

#### Time and setting
The experiments were performed at the Central Laboratory of Third Xiangya Hospital, Central South University, China from October 2009 to October 2010.

#### Materials
A total of 20 Wistar neonatal rats, within 1–3 days of birth, were provided by the Department of Experimental Animals, Xiangya Medical College, Central South University, China [license No. SCXK (Xiang) 2006-0002]. All protocols were conducted in accordance with the *Guidance Suggestions for the Care and Use of Laboratory Animals*, formulated by the Ministry of Science and Technology of China\(^{37}\).

#### Methods
**Isolation and primary culture of retinal neurons**
The neonatal rats were sacrificed by drowning in 75% alcohol. After disinfection for 5 minutes, the eyeballs were provided by the Department of Experimental Animals, Xiangya Medical College, Central South University, China [license No. SCXK (Xiang) 2006-0002]. All protocols were conducted in accordance with the *Guidance Suggestions for the Care and Use of Laboratory Animals*, formulated by the Ministry of Science and Technology of China\(^{37}\).
were harvested in a sterile manner, and washed twice in D-Hanks solution. The eyeballs were cut open at 0.5 mm posterior to corneal limbus using a dissecting microscope (Olympus, Tokyo, Japan). The retina was dissociated in D-Hanks solution and cut into pieces, triturated into a single cell suspension, and digested with 0.05% trypsin (Gibco, Carlsbad, CA, USA) at 37°C for 15 minutes, which was terminated by Dulbecco’s modified Eagle’s medium/F-12 culture solution (Gibco) containing fetal bovine serum. The suspension was filtered through a stainless steel mesh, centrifuged at 800–1 000 r/min for 5 minutes, resuspended in medium, and centrifuged. This process was repeated twice. The upper suspension was harvested, and the number of cells was quantified using a light microscope (Olympus). Cells at a density of 1 × 10⁶/mL were added to 12 mm × 12 mm coverslips and incubated in polylysine-coated 6-well culture plates with 80% Dulbecco’s modified Eagle’s medium/F-12 in 5% CO₂ at 37°C. Maintenance medium 1 [Neurobasal (Gibco), B27 supplement, 0.06 g/L glutamine, 100 U/mL penicillin, and 50 μg/mL streptomycin] was used after 24 hours, and maintenance medium 2 (Neurobasal, B27 supplement, 100 U/mL penicillin, and 50 μg/mL streptomycin) was used at 72 hours. One third of the medium was renewed every 1–2 days. Cell morphology was observed daily using an inverted microscope (Olympus) to identify quantity and length of processes and cell adherence.

Identification of retinal neurons
For Nissl staining, cell slices of 5–6 day primary cultures of retinal neurons were fixed in 4% paraformaldehyde for 25 minutes, washed with double distilled water three times, 5 minutes each, mixed with 10 g/L toluidine blue solution (Invitrogen, Carlsbad, CA, USA), incubated at 54°C for 30 minutes, cooled, and washed twice with double distilled water with a wash of 5 minutes each, rapidly washed with 95% alcohol, dehydrated with absolute alcohol, cleared with xylene, and mounted in neutral gum. Five high power fields (400 x) from each coverslip were randomly observed, and the number of positive neurons in every 100 cells in every field of view was quantified using a light microscope. The positive neuron amount in each field of view was calculated using the formula: percentage of neurons (%) = number of neurons/total number of cells × 100%\(^{(1)}\). Measurements were conducted in triplicate, and the mean value was calculated.

For immunocytochemical staining, retinal neuronal and glial cells were observed using anti-neurofilament and anti-glial fibrillary acidic protein immunocytochemical staining methods\(^{(2)}\). Briefly, cells were harvested from the coverslips, fixed with 4% paraformaldehyde at room temperature for 30 minutes, washed with PBS three times, 3 minutes each, blocked with 0.3% methanol in H₂O₂ at room temperature for 20 minutes, incubated with PBS containing 1% fetal bovine serum and 0.5% Triton X-100 for 20 minutes, followed by PBS containing 5–10% normal goat serum for 20 minutes. The cells were incubated with rabbit anti-rat neurofilament (1:100; Maixin-Bio, Fujian, China) and glial fibrillary acidic protein (1:50; Sigma, St. Louis, MO, USA) monoclonal antibody at room temperature for 2 hours, then overnight at 4°C, then washed with PBS three times, for 3 minutes each. The negative control was treated with PBS rather than primary antibody. Cells were incubated with biotinylated goat anti-rabbit IgG (1:200; Sigma) and peroxide-labeled streptavidin (Gibco), followed by coloration with 0.01 M Tris-HCl (pH 7.4) containing 0.05% diaminobenzidine and 0.03% H₂O₂. Coloration time was controlled using a microscope to terminate the reaction time. Cells were dehydrated and photographed. Glial cells were identified as follows: after nuclei were hematoxylin counterstained for 25 seconds, cells were washed with tap water, dehydrated with gradient alcohol, cleared with xylene, and mounted with neutral gum. Cells stained brownish-yellow were neuron-positive staining, cells with brownish-yellow or brown stained particles in the cytoplasm and membrane were glial cell-positive staining. Five high power fields (400 x) from each coverslip were randomly observed, and the number of neurofilament and glial fibrillary acidic protein positive neurons in every 100 cells in every field of view was quantified. The percentage of positive cells out of the total number of cells was calculated. Measurements were conducted in triplicate, and the mean value was calculated.

Establishment of a retinal neuron model of high glucose
According to a previously described method, cells were treated with 0 mM (normal control), 5.5, 15, 25, and 35 mM glucose (Sigma) for 24, 48, and 72 hours. Cell survival rate was determined using the MTT assay, and the apoptosis rate detected was determined using flow cytometry. The optimal glucose concentration and intervention duration for establishing the high glucose model were confirmed according to experimental results.

MTT assay for retinal neuron survival
Primary cultures of retinal neurons for 3–4 days were trypsinized (0.25%). A single cell suspension was prepared using serum-free culture medium, and cells at a
concentration of 1–3 × 10⁵ cells/mL were seeded into 96-well culture plates, with 100 μL in each well. The culture media containing different concentrations of glucose was replaced at a cell confluency of 80%. After culture for 24, 48, and 72 hours, 20 μL MTT solution (0.5 mg/mL, Sigma) was added to each well and cultured for 4 hours, treated with 100 μL dimethyl sulfoxide after the supernatant was discarded, and shaken at room temperature for 10 minutes to completely dissolve crystals. Absorbance at 570 nm was detected using an enzyme-linked immunosorbent assay reader (Perkin Elmer, Turku, Finland). The experiment was conducted in triplicate, and the mean value was calculated. Higher absorbance value represented more surviving cells.

**Flow cytometry for apoptosis**

Cells treated with different concentrations of glucose for 24, 48, and 72 hours were harvested, and digested with 0.25% trypsin (200 μL) in each well. The reaction was terminated by adding 10% fetal bovine serum after the cells shrunk. Cells were detached by trituration, centrifuged at 1 000 r/min for 5 minutes, washed with PBS twice. Cells were resuspended using 250 μL binding buffer, and cell concentration was adjusted to 1 × 10⁶ cells/mL. The cell suspension (100 μL) was placed in a 5 μL flow tube, incubated with 1 μL annexin/fluorescein isothiocyanate (Bender MedSystems, Vienna, Austria) and 5 μL propidium iodide (20 μg/mL) for 15 minutes in the dark, followed by 400 μL PBS. Cell apoptosis was analyzed using flow cytometry (Becton Dickinson, San Jose, CA, USA).

**Statistical analysis**

Data were expressed as mean ± SD and analyzed using SPSS 13.0 software (SPSS, Chicago, IL, USA). Comparison of the multiple group mean was conducted using one-way analysis of variance, and multiple comparisons among groups were analyzed using the Least Significant Difference test. A value of \( P < 0.05 \) was considered statistically significant.

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**Author contributions:** Yu Liu conceived and designed the study, was responsible for funds, performed statistical analysis, and wrote and revised the manuscript. Xueliang Xu supervised and provided important assessment of study design, as well as administrative support. Renhong Tang, Guoping Chen, Xiang Lei, Limo Gao, Wenjie Li and Yu Chen designed the study, conducted experiments and analyzed data. All authors approved the final version of the paper.

**Conflicts of interest:** None declared.

**Ethical approval:** This study received permission from the Animal Ethics Committee of Central South University, China.

**Author statements:** The manuscript is original, has not been submitted to or is not under consideration by another publication, has not been previously published in any language or any form, including electronic, and contains no disclosure of confidential information or authorship/patent application/funding source disquisitions.

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