Apigenin-7-O-β-D-(−6''-p-coumaroyl)-Glucopyranoside Treatment Elicits Neuroprotective Effect against Experimental Ischemic Stroke

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Abstract

Stroke is the major cause of permanent disability and mortality in China. Apigenin-7-O-β-D-(−6''-p-coumaroyl)-glucopyranoside (APG) is a glycoside subtype of apigenin and has the antioxidant activity; however, whether and how it plays a neuroprotective role following cerebral ischemia remains unknown. In present study, we adopted the oxygen glucose/reperfusion model in PC12 cells, bilateral common carotid artery occlusion model in C57B6 mice and middle cerebral artery occlusion model in SD rats to observe the therapeutic effects of APG on ischemic stroke. We also discussed the underlying mechanism. Treatment with 0.4 μg/ml or 0.8 μg/ml APG promoted cell viability and proliferation, reduced LDH release and apoptotic cell death levels in PC12 cells. Treatment with 50 mg/kg or 100 mg/kg APG at 30 minutes after reperfusion improved neurological outcomes in vivo, as demonstrated by elevation of neurological scores in both mice and rats. It also increased the number of survival neurons in mice and reduced infarct volume in rats. APG also increased the contents of Mn-SOD and the phosphorylation level of STAT3, elevated the antioxidant activity and reduced oxidative productions. These findings revealed a neuroprotective effect of APG, which possibly induced by the STAT3 phosphorylation-mediated Mn-SOD up-regulation.

Key words: Apigenin-7-O-β-D-(−6''-p-coumaroyl)-glucopyranoside (APG); cerebral ischemia/reperfusion; oxidative stress; manganese-superoxide dismutase

Introduction

Stroke, especially the ischemic stroke, is still one of the leading causes of morbidity and mortality in China (1). Unfortunately, few medications or therapies have been identified to treat ischemic stroke in the clinic. The current FDA approved stroke therapeutic medication is only the tissue plasminogen activator (tPA). However, owing to its narrow therapeutic window (<4.5h) and safety concerns, only less than 5% of patients can benefit from this medication (2). Therefore, more novel maneuvers need to be revealed for stroke treatment.

It is generally accepted that multiple cellular events, such as ionic imbalance, excitotoxicity, oxidative stress, apoptosis, inflammation, and necrosis involve the pathogenesis of ischemic stroke injury (3). Accumulated evidences have shown that oxidative stress plays a significant role in the pathophysiology of cerebral ischemic/reperfusion injury. During cerebral ischemia/reperfusion, the overproduction of free radicals damages the neurons and glia cells; and in-
teracts with cellular biomolecules, such as nucleic acid, lipids, and protein (4). Additionally, the sustained of the oxidative stress mediates the cascade amplification of impinges including mitochondrial dysfunction, inflammation response, and mitochondria/PS3-dependent cell death pathway activation (5). Despite the crucial role of oxidative stress in the pathophysiological process of cerebral ischemic damage, and the promising neuroprotective results of antioxidants supplementation demonstrated in animal studies, the treatment with exogenous antioxidants failed to verify their favorable outcomes in clinical trials (6). Actually, there exerts a coordinated self-protective mechanism in brain itself, which including endogenous antioxidant capacity. Exploring a new medication or therapy that triggers at activating endogenous antioxidant system may provide a new avenue for development of neuroprotective approaches.

In searching for stroke therapeutic medications, many traditional Chinese herbs, and their active components have been demonstrated to exert neuroprotective effect against experimental ischemic stroke injury (7-10). As a member of Ranunculaceae family, Clematis tangutica (Maxim.) Korsh (Ranunculaceae) is widely distributed in southwest and northwest areas of China (11). In traditional Tibetan medicine (TTM), the whole plant was used to treat indigestion and to invigorate circulation (12). Apigenin-7-O-β-D-(6''-p-coumaroyl)-glucopyranoside (APG) is one of the major active component in Clematistangutica which was demonstrated by our group (13). The free radical scavenging activity of APG has been identified, and the antioxidant capacity of this glycoside flavone is predominantly attributed to its reodox properties (14). Apigenin has been identified to induce significant neuroprotective effect against Parkinson disease(15), Alzheimer’s disease (16,17) and ischemic stroke injury (18) in a variety of in vitro or in vivo experimental models. As a glycoside subtype of apigenin, whether APG confers neuroprotective effect against cerebral ischemia is still unknown. Moreover, the underlying mechanism of the neuroprotective effect induced by this component also needs to be clarified.

In the present study, we tried to determine the neuroprotective effect of APG in three different experimental stroke models in vitro and in vivo. These models include oxygen and glucose deprivation and reperfusion (OGD-R) in PC12 cell line, bilateral common carotid artery occlusion (BCCAO) in mice and middle cerebral artery occlusion (MCAO) in rats. We also examined whether the up-regulation of endogenous antioxidant enzymes mediates the neuroprotective effect induced by APG treatment.

### Materials and Methods

#### Materials

APG was isolated from *C. tangutica*, and the molecular formula was established as C$_{30}$H$_{26}$O$_{12}$ (molecular weight: 578). The chemical structure of this compound is shown in Supplemental Fig. 1. Purified APG (Purity > 98%) used in current study was supplied by the Pharmacy Department of Xijing Hospital, the Fourth Military Medical University, Xi’an, China.

Male Sprague Dawley (SD) rats aged 8-12 weeks and weighting 250-280g were purchased from The Experimental Animal Centre of the Forth Military Medical University. Male C57Bl mice aged 8-12 weeks and weighting 20-25g were also obtained from this Centre. All experimental procedures were carried out according to protocols approved by the Ethics Committee for Animal Experimentation of the Fourth Military Medical University (Xi’an, Shaanxi, China) and in accordance with the National Institutes of Health Guide for the Care and Use of Laboratory Animals.

Antibodies used in the current study were: anti-phosphorylated STAT3 (705Y) rabbit monoclonal antibody (Cell signal technology; 1:1000); anti-total STAT3 rabbit monoclonal antibody (Cell signal technology; 1:1000); anti-Mn-SOD rabbit polyclonal antibody (Millipore; 1:500); and GAPDH monoclonal antibody (Abcam; 1:2000), β-tubulin monoclonal antibody (Abcam; 1:1000).

#### In vitro Oxygen and Glucose deprivation and re-oxygenation (OGD-R) model and APG treatment

The PC12 cells were plated at a density of 3×10$^5$ cells per well in 6-well multi well or ×10$^5$ cells per well in 96-well multi well at 37°C in 5% CO$_2$ and 95% air in DMEM administrated with 10% fetal bovine serum, streptomycin (100 µg/ml) and penicillin (100 units/ml). OGD-R model was established as described previously (19). Briefly, PC12 cells were switched to d-Hanks buffer (OGD medium) and then to a modular incubator chamber subsequently. The chamber was flushed with a gas mixture of 95% N$_2$ /5% CO$_2$ for 30 minutes at room temperature at the rate of 3 L/min. After flushing, the chamber was sealed and placed in a 37°C thermostat container. OGD was carried out for 4h. Following OGD, the cells were incubated with dulbecco's modified eagle medium (DMEM, without fetal bovine serum) with the vehicle or different concentrations of APG for additional 20 h reperfusion under normal conditions. The concentrations of APG were 0.2 µg/ml, 0.4 µg/ml, and 0.8 µg/ml and dimethyl sulfoxide (DMSO) was
used as vehicle in this experiment.

**In vitro cell viability assay**

The cell viability was determined by 3-(4,5-dimethyl-2-thiazolyl)-2,5-diphenyl-2-H-tetrazolium bromide (MTT) cell viability examination. Briefly, cells were cultured at $15 \times 10^3$ cells per well in 96-well tissue plates. At the end of the culture period, cells were washed with phosphate-buffered saline (PBS) and MTT was added to each well for incubation at 37 °C for 4 h according to the manufacturer’s instructions. Then, the medium was replaced with 150 μl of DMSO. The optical density (OD) was recorded on a Universal Microplate Reader (Elx 800, Bio-TEK Instruments Inc., USA) at 490 nm. Cell viability was shown as percentage changes in the control value.

**Lactate Dehydrogenase (LDH) release assessment**

Cell cytotoxicity was quantitatively assessed by evaluating the activity of LDH that was released from the injured cells into the culture medium after OGD-R. Then the cells were treated with 0.5 % Triton X-100, and the media containing detached cells were collected and centrifuged subsequently to yield supernatant for the assay of LDH activity. The LDH activity was measured using an assay kit according to the manufacturer’s recommendations. LDH leakage was expressed as the percentage of the total LDH activity (LDH in the both medium and cells), according to the equation, LDH released (%) = (LDH activity in the medium/total LDH activity) × 100%. Cultures without OGD-R treatment (sham group) were represented as basal LDH release.

**Flow cytometric analysis**

Cell apoptosis was analyzed by flow cytometry (20). Briefly, the cells were washed with $1 \times$ annexin V-FITC binding buffer and then stained with annexin V-FITC and PI for 15 min at room temperature in a dark room. Subsequently, the stained cells were analyzed by flow cytometric method. Apoptotic and necrotic cells were quantitated by annexin V binding and PI uptake. The cell populations of annexin V-FITC+/PI– were calculated to represent apoptotic cells.

**Cell cycle analysis**

Cell cycle was evaluated by flow cytometry as previously described (21). The cells were collected by trypsinization and centrifuged by PBS twice, then fixed in pre-cooling 70% ethanol at -20 °C and subsequently stained with PI solution. DNA content was measured by flow cytometry using Cell Quest Software. Totally, 10,000 events were counted for each sample (FACSC alibur, Becton–Dickinson, USA). The percentage of cells in any particular cell cycle stage was calculated using the ModFit software (Becton–Dickinson, USA).

**Global cerebral ischemia/reperfusion model establishment in mice**

In the current study, global cerebral ischemia/reperfusion was induced by BCCAO operation. Mice were inhaled with 3 % isoflurane for anesthetic induction, and the anesthetic was maintained with 1.5 % isoflurane inhalation. After a midline incision between the neck and sternum had been made, both common carotid arteries were located lateral to the sternocleidomastoid. The surrounding tissues and vagus nerve were separated carefully. Global cerebral ischemia was induced by clamping the common carotid arteries with two miniature artery clips. After 20 min of ischemia, the clips were removed to allow the blood reflow. During the surgical procedure, the pericranial temperature was maintained at 37.0 ± 0.5 °C with a heating pad. Additionally, regional cerebral blood flow (rCBF) was monitored by a laser Doppler flowmeter (PeriFlux System 5000; Perimed, Stockholm, Sweden). The measurement started from the time of anesthetic induction till 5 minutes after reperfusion. In this experiment, only mice whose cortical rCBF was suppressed to <15% of baseline during ischemia and was recovered to 85% during reperfusion process were included for data analysis. Mice were treated with the vehicle or different concentrations of APG via intraperitoneal injection at 30 minutes after reperfusion. The concentrations of APG were 25 mg/ml, 50 mg/ml, and 100mg/ml and 10% DMSO was used as vehicle in this experiment.

**Neurobehavioral evaluation in mice**

At 72 h after reperfusion, mice were subjected to modify neurological measurements designed to detect motor deficits by an observer unaware of the grouping. Tree tests were performed in this experiment. In the first test, a 10×20 cm screen with 0.2cm×0.2cm grid size, which could be rotated from horizontal to vertical, was used. During the test, the mouse was placed on the horizontal screen, and then the screen was rotated into a vertical plane. The duration time that the mouse held onto the screen was recorded for a maximum of 15 s. Next, the mouse was put in the middle of a horizontal wooden rod with a diameter of 1.5 cm. The time that the mouse was able to remain balanced on the rod was recorded for a maximum of 30 seconds. Finally, a prehensile traction test was performed. The time that the mouse could cling to a horizontal rope was recorded for a maximum of 5 seconds. Every test was allowed a total of 3 points. From 3 tests, a total motor score was computed for 9
possible points.

**Histological evaluation of neuronal damage in mice**

The evaluation of histological damage was performed 72 hours after reperfusion when the neurological test had finished. After perfused with freshly ice-cold paraformaldehyde in 0.01 M phosphate buffered saline (PBS), brains were removed, and coronal sectioned. Then sections that included the dorsal hippocampus were stained with hematoxylin-eosin, thus the pyramidal neurons in CA1 region could be stained. The total number of viable neurons in the CA1 region was counted in a blind manner by light microscopy at ×400 magnification in 3 different fields for each section. Data from 6 mice in each group were averaged.

**Transient focal cerebral ischemia/reperfusion model in rats**

Middle cerebral artery occlusion model was used to induce transient focal cerebral ischemia reperfusion injury as described previously (22). For the preparation of surgery, the rats were fasted overnight with free access to tap water. Rats were anesthetized with 10% chloral hydrate (350 mg/kg). To obstruct blood flow in MCA, a 3-0 nylon suture with blunted head was inserted into internal carotid artery through the right common carotid artery until a middle resistance was felt. The occlusion was maintained in position for 2 h and subsequently removed to allow blood reperfusion. The administration of APG was confirmed according to the paradigm used in mice, as described above.

**Neurological score and infarct volume evaluation in rats**

The neurological outcome was evaluated at 72 h after reperfusion by a blinded observer according to Garcia’s methods.

After neurological outcome examination, rats’ brains were removed and sectioned. The infarct volume was evaluated as described previously. Briefly, the slices were stained with 2% solution of 2,3,5-triphenyltetrazolium chloride at 37 °C and then transferred to 4% paraformaldehyde for 24 h. The brain slices were photographed using a digital camera (Canon Ixus 220HS, Japan). The infarct volume was calculated according to the following equation: relative infarct size = (contralateral area - ipsilateral non-infarct area) / contralateral area.

**Western blot assessment**

Tissue from MCAO models was used to investigate the mechanism of APG induced neuroprotective effect. The penumbra tissue of the anesthetized rats from another independent groups was promptly removed at 2 h after reperfusion (n = 5 per group). Tissue was homogenized in ice-cold RIPA lysis buffer (Beyotime, Nantong, China) mixed with 1% phenylmethanesulfonyl fluoride (PMSF). Equal amount of protein (40 μg) was loaded into each lane of a polyacrylamide-SDS gel and electrophoresed. The resolved proteins were transferred to a PVDF membrane. Membranes were blocked with 5% bovine serum albumin (BSA) and incubated with the appropriate primary antibodies overnight at 4 °C. The membranes were each washed three times in Tris-Buffered Saline Tween-20 (TBST) for 5 min. Then the membranes were incubated for 1 h at room temperature with the appropriate secondary antibodies. The immune-blots were immersed in enhanced chemiluminescent (ECL; Millipore) reagent to analyze specific protein. Protein expression or activation changes were presented with the ratio of β-tubulin or GAPDH.

**ELISA measurement of antioxidants and oxidative productions**

Penumbra samples from the ischemic hemisphere were harvested and homogenized in ice-cold saline with a 1:10 weight-to-volume value. Then the samples were centrifuged at 3000 rpm for 15 min at 4 °C and the supernatant was collected for ELISA assessment. The contents of antioxidant enzymes, total superoxide dismutase (T-SOD, Beyotime), catalase (CAT, Beyotime) and total antioxidant capacity (T-AOC, Beyotime); the oxidative productions, malondialdehyde (MDA, Beyotime), protein nitrotyrosine (Millipore) and reactive oxygen species (ROS, West-tone) were measured by available ELISA kits, according to their instructions.

**Statistical Analysis**

Nonparametric values such as neurological scores were analyzed by a Kruskal-Wallis test followed by the Mann-Whitney U statistic with Bonferroni correction. Other values were presented as mean ± SD and were analyzed by one-way analysis of variance, followed by Bonferroni correction for post-hoc T-test. Values of P<0.05 were considered statistically significant.

**Results**

**Protective effect of APG against OGD-R injury**

As shown in Figure 1, OGD-re-oxygenation reduced cell viability to 64±5.13% of the sham group and APG at administration of 0.2 μg/ml showed no significant effect on increasing the cell viability. Treatment with APG at 0.4 μg/ml and 0.8 μg/ml restored the cell viability (Fig. 1A, P<0.05 vs. the control
group, respectively). However, no dose-dependent effect was detected in these two groups.

Compared to the sham group, the LDH level in the cells of OGD-R groups was increased significantly. Treatment with APG at 0.2 ug/ml did not reduce the elevated LDH level induced by OGD-R injury. APG treatment at 0.4 ug/ml and 0.8 ug/ml showed significant protective effect on attenuation of the increased LDH level induced by OGD-R, as compared to OGD group (Fig. 1B, 53.4% reduction in 0.4 ug/ml group, P<0.01; 45.3% reduction in 0.8 ug/ml group; P<0.05).

To test the effect of APG treatment on cell apoptosis, flow cytometric analysis was also conducted in this study (Fig. 2A-E). The apoptotic index of the control and vehicle group were increased in comparison with the sham group (Fig. 2K). Both treatment with 0.4 ug/ml and 0.8 ug/ml APG attenuated OGD-R-induced cell apoptosis significantly as compared with that in OGD-R group (63.8% reduction in 0.4 ug/ml group; 59.2% reduction in 0.8 ug/ml group; P<0.01, respectively). Conversely, treatment with 0.2 ug/ml APG had no protective effect on cell apoptosis.

Figure 1. The protective effect of APG against OGD-R injury in PC12 cell line. Effect of APG treatment on the cell viability (A) and plasma lactate dehydrogenase (LDH) releasing level (B) in the PC12 cell culture after OGD-R. A, cell viability analysis; B, LDH releasing level evaluation. APG treatment significantly increased cell viability and decreased the LDH releasing level compared with OGD group. *P < 0.05 vs. sham group; #, ##P < 0.05, P < 0.01 vs. OGD group. n = 5 per group.

Figure 2. Effect of APG therapy on the apoptotic level and cell proliferation in PC12 cells. A-E. Representative dot plots showing flow cytometric analysis of PC12 cells treated with different concentrations of APG for the dedicated time periods and then stained with FITC-conjugated Annexin V and propidium iodide in OGD (A), vehicle (B) and 0.2 ug/ml (C), 0.4 ug/ml (D), 0.8 ug/ml (E) APG treatment groups, respectively. F-J. Flow cytometric analysis of cell cycle phase distribution of PC12 cells after treatment with different concentrations of APG or vehicle (F, control; G, vehicle; H, 0.2 ug/ml; I, 0.4 ug/ml; J, 0.8 ug/ml). K. Evaluation of apoptotic index in each group (n = 5). L. Analysis of proliferation index in each group (n = 5). All concentrations of APG administration increased the percentage of S+G2 cells but not altered the percentage of S cells (the left legend is the percentage of S cells, the left legend is the percentage of S+G2 cells). *P < 0.05, **P < 0.01 vs. sham group.
The effect of APG treatment on cell proliferation was also measured in the present study (Fig. 2F-J). Treatment of PC12 cells with different doses of APG increased the percentage of the S and G2 phase cells as compared with sham group, however no dose-dependent changes were found (Fig. 2I, 38.2%, 50.9%, 45.5% improvement in 0.2 μg/ml group, 0.4 μg/ml group, 0.8 μg/ml group; P<0.01, respectively).

**Neuroprotective effect of APG treatment in a global cerebral ischemia/perfusion mice model**

As shown in Fig. 3A, there was no significant difference in rCBF changes among all groups at the corresponding time points (Fig. 3B). No mice died until the neurological assessment at 72 h after reperfusion. In the BCCAO and vehicle groups, the total motor scores were reduced in global ischemia/reperfusion group, as compared with the one in sham groups. Both treatment with 50mg/kg (7.67[7-9]) and 100 mg/kg (7.67[7-9]) APG attenuated the total motor scores deficiency evidently (Fig. 3B, in 100mg/kg versus 5.9[5-7] in BCCAO group). However, no statistical difference was detected between these two different dose groups. No protective effect was found in 25 mg/kg APG treatment group (6.8 [6-8]).

Figure 3. APG treatment improved the total motor scores (B) and protected hippocampal CA1 neurons (C) against global ischemic/reperfusion injury. A. Changes in regional cerebral blood flow (rCBF) in C57BL/6 mice during BCCAO operation. Data are expressed as mean ± SD. B. Triangle, square and circle symbols represent mice in each group (n = 12), respectively. C. Counting of viable neuronal cells in hippocampal area in each group (n = 12). D. Representative microphotographs of hematoxylin eosin staining in sham, BCCAO, vehicle, 25 mg/kg, 50 mg/kg and 100 mg/kg APG treatment groups. * P < 0.05 vs. sham group; # P < 0.05, ## P < 0.01 vs. BCCAO group. BCCAO = bilateral common carotid artery occlusion. Bar = 20 μm.
The histopathological results are shown in Fig. 3C-D. Viable neurons in hippocampal CA1 region were significantly decreased at 72 h after ischemia/reperfusion compared with which in the sham group. Treatment with 50 mg/kg or 100 mg/kg APG increased the number of reduced viable neurons induced by ischemia/reperfusion injury as compared with the BCCAO group. However, administration with 25 mg/kg APG had no protective effect on viable neurons.

**Neuroprotective effect of APG in a rat focal cerebral ischemia/perfusion model**

No significant difference in rCBF changes was detected among all groups at the corresponding time points (Fig. 4A). The neurological score assessment is presented in Fig. 4B. The MCAO group revealed a significant neurological deficit score when compared to the sham group. Treatment with APG significantly improved the neurological scores, in 50mg/kg (13.63±10-16) or 100mg/kg (12.38±9, 14) respectively, as compared with that in the MCAO group (8.0±6-12). However, no statistical difference was detected in neurological score between the MCAO group and 25 mg/kg APG treatment group (9.62±7-13).

As shown in Fig. 4C-D, cerebral I/R injury induced a 41.12% infarction volume in the MCAO group while the sham group showed no infarction area. Compared with the control group, the cerebral infarct volumes had a 30.0% deduction in 50mg/kg APG group (P < 0.01) and a 23.3% deduction in 100mg/kg APG group (P<0.05). No difference was detected between the latter two groups.

**APG treatment promoted STAT3 phosphorylation**

As shown in Fig. 5A, the difference of STAT3 phosphorylation level between MCAO and vehicle group was not statistically significant. APG treatment increased the phosphorylation of STAT3 at 705Y as compared to the MCAO group at 2 h after reperfusion (P<0.05). However, no significant change was detected in overall expression of total STAT3 among all the groups.
Figure 5. Effect of APG therapy on the phosphorylation of STAT3 at 705Y (A) and the expression of Mn-SOD (B). A. The upper part is the photograph of pSTAT3 (705Y), total STAT3, and the corresponding GAPDH bands, respectively. The lower part is the result of western-blot statistical analysis in the sham, control, vehicle and APG treatment groups (n = 5). APG treatment increased STAT3 phosphorylation as compared with the MCAO group. B. The upper part is the photograph of Mn-SOD and its corresponding β-tubulin bands. The lower part is the result of western-blot statistical analysis in the sham, MCAO, vehicle and APG treatment groups (n = 5). APG treatment increased Mn-SOD2 expression as compared with the control group. * P < 0.05 vs. sham group; # P < 0.05 vs. MCAO group. MCAO = middle cerebral artery occlusion.

APG treatment upregulated expression of Mn-SOD protein

As shown in Fig. 5B, ischemic/reperfusion injury attenuated the content of SOD2 protein as compared with that in the sham group. There is no statistical significant difference in Mn-SOD protein expression between MCAO and vehicle group subjected to MCAO injury. APG treatment increased the expression of Mn-SOD as compared with the control group at 2 h after reperfusion (P<0.05).

APG therapy increased protective antioxidant effect

The impact of APG treatment on antioxidant enzymes expression was compared with the MCAO groups at 24 h after reperfusion. Some antioxidant enzymes, CAT, T-AOC, and T-SOD were activated by APG treatment as measured by ELISA kits (Fig. 6A-C; P<0.05, respectively).

Some oxidative productions were also measured in the current study. As shown in Fig. 6D-E, the content of MDA, nitrotyrosine and ROS increased in the MCAO group compared to the sham group. APG treatment attenuated the expression of MDA and ROS compared to the MCAO group (P<0.05, respectively).

Discussion

In present study, we demonstrated for the first time as we known that APG exhibited neuroprotective effects against cerebral ischemic/reperfusion injury. APG markedly increased cell proliferation ratio, reduced the number of apoptotic cells; improved cell viability and attenuated LDH releasing level against OGD-R as demonstrated in vitro. Also, APG treatment significantly ameliorated neurological behavior, elevated survival neurons and reduced infarct volume in both global cerebral ischemic and focal cerebral ischemic models in vivo. Furthermore, the results of the current study showed that APG treatment dramatically reduced levels of MDA, protein nitrotyrosine, ROS, as well as improved the content of antioxidant enzymes. In addition, APG treatment increased Mn-SOD protein expression and the phosphorylation level of STAT3 at 705Y. These results indicate that the antioxidant activity induced by APG treatment was mediated by Mn-SOD up-regulation via STAT3 phosphorylation dependent mechanism, which may be one of the probable mechanisms of the neuroprotection induced by APG treatment.

Accumulating studies suggest that some bioactive compounds discovered from traditional Chinese medicines such as apigenin can attenuate cerebral ischemic/reperfusion injury in the animal models (23). As a glycoside flavonoid, APG is one of the major compounds isolated from C. tangutica. The molecular weight of APG is small than 3,000, and it is a liposoluble constituent. Therefore, APG is believed to easily and rapidly cross the blood-brain barrier and exert biological activities in the brain.
Currently, we have demonstrated that APG can confer neuroprotection in three considerable ischemic/reperfusion injury models, respectively. In PC12 cell line, APG improved cell viability, increased cell proliferation ratio, attenuated LDH releasing level and reduced the number of apoptotic cells. Additionally, we have also shown that APG can induce neuroprotective effect in global cerebral ischemic/reperfusion model of mice, as evidenced by improved neurological function and increased survival neurons. Moreover, APG also ameliorated neurological function and attenuated infarct volume measurement in a transient rat MCAO model. The neuroprotective dose of APG at 0.4 μg/ml in vitro or 50 mg/kg in vivo was more potent than those at 0.2 μg/ml in vitro or 25 mg/kg in vivo. However, the higher dose of APG (0.8 μg/ml in vitro or 100 mg/kg in vivo) did not show any more protective potency, indicating that the middle dose (0.4 mg/ml in vitro or 50 mg/kg in vivo) of APG might be an appropriate dose for the further studies.

It is well established that the overproduction of ROS is a crucial injurious mechanism in the context of I/R injury (24). During I/R process, suddenly accumulated ROS cannot be efficiently scavenged by the endogenous antioxidant enzymes, which subsequently interacts with subcellular biomolecules, causing the cascade amplification of injury pathogenesis such as mitochondrial dysfunction, apoptosis (24,25). Therefore, raising the expression of endogenous antioxidant enzymes can efficaciously reduce cerebral I/R injury. The prominent free radical scavenging activity of APG has been identified (26, 27). In the present study, we have demonstrated that APG increased the expression of antioxidant enzymes including SOD, CAT, and T-AOC. These enzymes constitute the first-line cellular defense system against oxidative injury, thus the activation of antioxidant enzymes induced by APG treatment could induce neuroprotective effect.

The half-life of ROS such as O$_2^-$ was very short. Thus, it is very difficult to real-time monitor them in vivo. The breaking out of ROS induced by I/R also induces the damage of biomolecules, such as lipid and protein. So the assessment of oxidative biomolecules was constantly used to evaluate the oxidative injury. In the current study, we demonstrated that APG reduced the nitrotyrosine protein, MDA and ROS levels in brain tissue. This result indicates the involvement of the antioxidant effect in the neuroprotection induced by APG. However, the underlying mechanism of the antioxidant effect induced by APG is still unknown.

Mn-SOD primarily mediated the mitochondrial
mechanism to eliminate or maintain the homeostatic level of $O_2^-$, to process ROS into fewer toxic substances (28, 29). It is also an important particle of endogenous protective mechanism against cerebral ischemic damage. In this study, we have shown that the Mn-SOD protein expression was suppressed in the penumbra at 2 h after reperfusion, which was in consistent with previous studies (30). In addition, in the APG-treated brain, the content of Mn-SOD protein was much more pronounced than that in mice subjected to only cerebral I/R injury. This result indicated the involvement of Mn-SOD in the neuroprotective effect induced by APG. The up-regulation of Mn-SOD can undergo transcriptional, translational and post-translational regulation mechanism, which induced protective effect against oxidative injury. However, it is still unclear which upstream signal pathway transduced the activation of Mn-SOD induced by APG. As a signaling transducer, the activation of STAT3 mediated much subcellular protective mechanism, also including the antioxidant effect (31, 32). It has been identified that STAT3 was a novel transcription factor of the Mn-SOD gene (30), in present study, we have demonstrated that APG increases the STAT3 phosphorylation level at 705Y, indicating that APG induces the Mn-SOD up-regulation via STAT3 phosphorylation dependent-mechanism and then attenuates the oxidative stress, resulting in neuroprotective effect.

Limitations
Some limitations should be pointed out in this study. Firstly, more experimental evidences are needed to determine the correlation of Mn-SOD up-regulation and the neuroprotection induced by APG. Moreover, as a transcriptional factor, the phosphorylation of STAT3 could be increased by APG, but more investigations are needed to determine the upstream pathway that transduced the phosphorylation of STAT3 induced by APG.

Conclusion
Our study provided both in vitro and in vivo evidence that APG exerted neuroprotective effects against cerebral ischemic/reperfusion injury. Although much more studies are still needed to explain the involvement of transduction signal pathways between APG and antioxidant system fully, the present study demonstrates that APG treatment can provide ischemic tolerance to cerebral ischemic oxidative injury. This was mediated by activation of Mn-SOD signaling pathway via STAT3 phosphorylation dependent-mechanism. Thus, these results indicate that APG may be a potent therapeutic drug for the treatment of ischemic stroke.

Supplementary Material
Supplementary Figure 1. http://www.ijbs.com/v12p0042s1.pdf

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Competing Interests
The authors have declared that no competing interest exists.

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