Gynura bicolor aqueous extract attenuated H2O2 induced injury in PC12 cells

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ABSTRACT

Background: Protective effects of Gynura bicolor aqueous extract (GAE) at three concentrations upon nerve growth factor (NGF) differentiated-PC12 cells against H2O2 induced injury were examined.

Methods: NGF differentiated-PC12 cells were treated with GAE at 0.25%, 0.5% or 1%. 100 μM H2O2 was used to treat cells with GAE pre-treatments. After incubating at 37°C for 12 hr, experimental analyses were processed.

Results: H2O2 exposure decreased cell viability, increased plasma membrane damage, suppressed Bcl-2 mRNA expression and enhanced Bax mRNA expression. GAE pre-treatments reversed these changes. H2O2 exposure reduced mitochondrial membrane potential, lowered Na+-K+-ATPase activity, and increased DNA fragmentation and Ca2+ release. GAE pre-treatments attenuated these alterations. H2O2 stimulated the production of reactive oxygen species (ROS), interleukin (IL)-1-beta, IL-6 and tumor necrosis factor-alpha, lowered glutathione content, and reduced glutathione peroxidase (GPX) and catalase activities. GAE pre-treatments maintained GPX and catalase activities; and concentration-dependently diminished the generation of ROS and inflammatory cytokines. H2O2 enhanced mRNA expression of nuclear factor kappa (NF-κ) B and p38. GAE pre-treatments decreased mRNA expression of NF-kB and p38. Conclusion: These findings suggested that GAE might be a potent neuronal protective agent.

Keywords: Gynura bicolor; NGF-PC12 cell; Apoptosis; NF-κB; p38

1. Introduction

Oxidative and inflammatory reactions are involved in nigral degeneration and neuronal cell death, which contribute to the pathogenesis of neurological disorders such as Parkinson’s disease (PD) [1]. The over-generated oxidants and inflammatory cytokines including reactive oxygen species (ROS), interleukin (IL)-1Β, IL-6 and tumor necrosis factor (TNF)-alpha cause neuronal cells apoptosis, impair brain functions, and deteriorate PD and/or other neurological diseases [2]. In addition, the increased caspase activity, Bax expression, and nuclear transcription factor kappa (NF-κ) B activation due to some stimuliants also promote damage, and even death of neuronal cells [3,4]. Thus, exploring the appropriate natural agent(s) with the capabilities to decrease the production, activity or expression of these above factors might be a good and safe choice in order to enhance the stability of neuronal cells, prevent or attenuate the progression of neurological disorders. PC12 cell line, a rat adrenal gland pheochromocytoma cell line, could become a sympathetic neuronal phenotype through reacting with nerve growth factor (NGF) for differentiation [5]. So far, NGF treated PC12 cells have been considered as sympathetic neurons to investigate the protective effects and action modes of some potent compounds for neuronal cells [6,7].

Gynura bicolor DC. (G. bicolor) is a plant food, and available in several Asian countries such as China, Taiwan, Japan and Malaysia. Its leaf part is an edible vegetable. G. bicolor has been applied in folk medicine for diabetes treatment in China southern area [8]. Tuekpe et al. [9] reported that dietary G. bicolor intake promoted urinary potassium excretion, which benefited the management of blood pressure for healthy Japanese women. The study of Teoh et al. [10] revealed that component compounds of G. bicolor exhibited cytotoxic effects for colon cancer cells. Wu et
al. [11] reported that G. bicolor water or ethanol extract enhanced iron bioavailability in rats. In the study by Chao et al. [12], four groups of phytochemicals including flavonoids, phenolic acids, carotenoids and anthocyanins were detected in aqueous extract of G. bicolor leaf part, and their content were 1934, 1428, 921 and 2135 mg/100 g dry weight. Furthermore, this aqueous extract displayed anti-oxidative activities for human umbilical veno-endothelial cells against high glucose [12]. In addition, our previous animal study found that dietary intake of G. bicolor aqueous extract (GAE) markedly attenuated hepatic glycative injury and lipid accumulation in mice with chronic ethanol consumption [13], and the authors indicated that the observed hepatic protective activities from GAE were due to the contribution of its phytochemical component compounds. These previous studies suggest that GAE could offer multiple bioactivities. Therefore, it is hypothesized that GAE might be able to protect neuronal cells.

In order to understand whether GAE could be developed as a neuro-protective agent, our present cell line study was conducted. NGF differentiated-PC12 cells were pre-treated with GAE at three concentrations. Then, hydrogen peroxide was used to induce apoptotic, oxidative and inflammatory stress. The effects of GAE on cell survival, plasma membrane integration, caspases and Na\(^+\)-K\(^+\)-ATPase activities, and mRNA expression of Bel-2, Bax, NF-sxB and p38 were examined. Furthermore, the anti-oxidative and anti-inflammatory activities of GAE against H\(_2\)O\(_2\) were also evaluated. These results could partially support and explain the possibility of considering GAE as a neuro-protective nutraceutical.

2. Materials and methods

2.1. Materials

Fresh G. bicolor was directly purchased from farms in spring, 2015. 100 gram fresh leaf part was cut into small pieces, and mixed with 250 ml double distilled water. After homogenizing in a blender, GAE was collected via filtering through a No. 1 whatman filter paper. GAE was further freeze-dried to fine powder. The content of total phenolic acids and total flavonoids in GAE were in the range of 1428 ± 137 and 1934 ± 108 mg/100 g fine powder [12]. In our present work, the levels of total phenolic acids and total flavonoids were measured in order to standardize the used GAE. NGF was purchased from Sigma Chemical Co. (St. Louis, MO, USA). Antibodies were obtained from Boehringer-Manheim Co. (Indianapolis, IN, USA). Culture medium, plates and chemicals for cell culture were bought from Difco Laboratory (Detroit, MI, USA).

2.2. PC12 cell culture and treatments

PC12 cells cultured in Dulbecco’s modified Eagle’s medium (DMEM) were routinely maintained under 95% air and 5% CO\(_2\) at 37°C. PC12 cells were treated by NGF at 50 ng/mL, and followed by a 5-day incubation at 37°C. Medium was refreshed every 72 hr. After washing twice with serum-free DMEM, cells were collected and loaded in 96 well plates. Cell number was adjusted to 10\(^3\)/ml by phosphate buffer saline (PBS). GAE was dissolved in DMEM. Two groups of NGF differentiated-PC12 cells were treated with 500 μL DMEM only; they were a normal group and a control group, respectively. Three groups of NGF differentiated-PC12 cells were treated with 500 μL DMEM containing GAE at 0.25%, 0.5% or 1%. After incubation for 48 hr at 37°C, cell samples were washed twice with serum-free DMEM. Then, those used serum-free DMEM was collected, and the content of phenolic acids, flavonoids, carotenoids or anthocyanins was analyzed according to the methods described in Chao et al. [12]. There were no detectable phenolic acids, flavonoids, carotenoids or anthocyanins in the DMEM used for washing. Subsequently, 100 μM H\(_2\)O\(_2\) was used to treat control group, and three groups of cells with GAE pre-treatments. After incubating at 37°C for 12 hr, experimental analyses were processed.

2.3. Cell survival and plasma membrane damage

Cell survival was measured by 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide (MTT) assay. In brief, MTT at 0.25 mg/mL was added into cell suspension, and this mixture was incubated at 37°C for 3 hr. MTT formazan product was quantified by monitoring the absorbance at 570 nm by a Bio-Rad microplate reader (Hercules, CA, USA). Cell viability was presented as a percentage of normal groups. Plasma membrane damage was assayed by determining lactate dehydrogenase (LDH) activity. After centrifugation, 50 μL supernatant was used to measure LDH activity (U/L) by a kit (Sigma Chemical Co., St. Louis, MO, USA) according to manufacturer’s instruction.

2.4. Assays for DNA fragmentation and mitochondrial membrane potential (MMP)

DNA fragmentation was determined by a cell death detection ELISA kit (Roche Molecular Biochemicals, Mannheim, Germany) according to manufacturer’s instruction. Cells were suspended in cold lysis buffer for 30 min at 25°C, and centrifuged for 10 min at 250 g. Twenty μL supernatant was used to react with 80 μL freshly prepared immunoreagent, and followed by incubating for 2 hr at 25°C. After washed twice with PBS, substrate was added and followed by incubating for 15 min at room temperature. A microplate reader was applied to monitor the absorbance at 405 nm and 490 nm. DNA fragmentation was shown as an enrichment factor, which means: (absorbance of the sample) / (absorbance of the control groups). MMP was measured by using Rh123, a fluorescent dye. Cell samples were treated with Rh123 at 100 μg/L for 30 min at 37°C. After washed twice with PBS, the mean fluorescence intensity (MFI) was analyzed by a Beckman-FC500 flow cytometry (Beckman Coulter, Fullerton, CA, USA).

2.5. Measurement of caspases and Na\(^+\)-K\(^+\)-ATPase activities

Caspase-3 and caspase-8 activities were quantified by fluorometric kits (Upstate, Lake Placid, NY, USA) according to manufacturer’s instructions. Cells were lysed, and protein concentration was determined by a Pierce assay kit (Rockford, IL, USA). The lysates were reacted with specific substrates, and followed by incubating 60 min at 37°C. Fluorescence value was recorded by a Hitachi F-4500 fluorophotometer (Tokyo, Japan), in which excitation and emission wavelengths were 400 nm and 505 nm. The variability coefficients of inter-assay and intra-assay were 3.9-5.6% and 4.3-5.9%, respectively. Caspase-3 or caspase-8 activity was defined as fluorescence unit/mg protein. Na\(^+\)-K\(^+\)-ATPase activity was assayed according to the method of Torlinska and Grochowska [14] via analyzing the released amount of inorganic phosphate (Pi) from ATP. The released Pi was determined
by monitoring the absorbance at 640 nm. The value of the treated groups was shown as a percentage of normal groups.

### 2.6. Assay of intracellular Ca\(^{2+}\) level

A Ca\(^{2+}\)-sensitive dye, Fura-2-AM, was used to detect the intracellular Ca\(^{2+}\) level via recording the change in fluorescent intensity [15]. In brief, Fura-2-AM at 5 mmol/L was added into cells (105 cells/mL), and stored in dark condition for 30 min at 25°C. After further incubating 30 min at 37°C, fluorescence value was recorded by a Shimadzu spectrofluorimeter (Model RF-5000, Kyoto, Japan). The emission wavelength was set at 510 nm, and excitation wavelength was set at 340 and 380 nm. Calcium concentration (nM) was calculated according to the equation: [Ca\(^{2+}\)] = Kd ∗ [(R–Rmin)/(Rmax–R)] ∗ FD/FS. Kd was 224 nM, R was the ratio of fluorescence values at 340 and 380, Rmax was measured by using triton X-100 to treat cells, Rmin was measured by using ethylene glycol tetraacetic acid to treat cells. FD was the fluorescence value of Ca\(^{2+}\)-free form, and FS was the fluorescence value of Ca\(^{2+}\)-bound form at 340 and 340 nm.

### 2.7. Assays for oxidative and inflammatory associated factors

ROS level was determined by 2',7'-Dichlorofluorescein diacetate (DCFH-DA). In brief, 100 μL cell homogenate was mixed with 100 μL 2 mg/mL DCFH-DA. After incubating at 37°C for 30 min, fluorescence value was recorded by a Gemini XS fluorescence plate reader (Molecular Devices, Sunnyvale, CA, USA). Emission and excitation wavelengths were 525 nm and 488 nm, respectively. Result was expressed as relative fluorescence unit (RFU) per mg protein. The level of glutathione (GSH), and the activity of glutathione peroxidase (GPX) or catalase were measured via assay kits purchased from OxisResearch Co. (Portland, OR, USA) according to manufacturer’s instructions. The levels (pg/mg protein) of IL-1beta, IL-6 and TNF-alpha were quantified by cytoscreen assay kits obtained from BioSource International (Camarillo, CA, USA). The detection limit was 5 pg/mg protein.

### 2.8. Real-time polymerase chain reaction (RT-PCR) for mRNA expression

Total mRNA of cells was extracted by reagents obtained from Invitrogen Trizol (Life Technologies, Carlsbad, CA, USA). RNA concentration was quantified by monitoring the absorbance at 260 nm. Subsequently, 5 μg RNA was applied for generating cDNA via reverse-transcription procedure. Then, cDNA was further used for PCR process. The primers of target genes were as follow: Bcl-2: forward, 5'-CGT TTG GCA GTG CAA TGG T-3'; reverse, 5'-TTC TTG ATT GAG CGA GCC TT-3'; Bax: forward, 5'-TGG CAG CTC ACA TGT TTT CTG AC-3'; reverse, 5'-TCA CCC AAC CAC CCT GGT CTT-3'; NF-kB, forward, 5'-GAG GTC TCT GGG GGT ACA GTC-3'; reverse, 5'-GGA CAA CGC AGT AGA ATT TTA-3'; p38: forward, 5'-GCC AAG GGC TAC ACC AAA TC-3'; reverse, 5'-GAC TCA TGA CCA CAG TCC ATG C-3'. PCR amplification condition was 3 min denaturation at 95°C, 10 s annealing at 60°C and 20 s extension at 72°C. For Bcl-2, Bax, NF-kB or p38, 35 cycles were performed; for GADPH, 28 cycles were processed. A Sequence Detection System (ABI Prism 7700, Applied Biosystems, Foster City, CA, USA) was used to quantify PCR products.

### 2.9. Statistical analyses

Data were obtained from 7 different preparations, and expressed as mean ± standard deviation (SD) (n = 7). Statistical analyses were processed by using one-way analysis of variance. Dunnett’s t-test was applied for Post-hoc comparison. P value lower than 0.05 was defined as significant.

### 3. Results

#### 3.1. GAE alleviated apoptosis and plasma membrane damage

Without H\(_2\)O\(_2\) stimulation, GAE at test concentrations did not affect viability and plasma membrane stability in NGF differentiated-PC12 cells (Fig. 1a and 1b, P < 0.05). As shown in Fig. 2, H\(_2\)O\(_2\) exposure decreased cell viability (2a) and increased plasma membrane damage (2b), determined by LDH activity, in NGF differentiated-PC12 cells when compared with normal groups (P < 0.05). GAE pre-treatments concentration-dependently increased cell viability and diminished LDH activity (P < 0.05). H\(_2\)O\(_2\) reduced Bcl-2 mRNA expression and enhanced Bax mRNA expression in NGF differentiated-PC12 cells (Fig. 3, P < 0.05). GAE pre-treatments at test concentrations raised Bcl-2 mRNA expression (P < 0.05); and concentration-dependently lowered Bax mRNA expression (P < 0.05).

#### 3.2. GAE attenuated mitochondrial and DNA injury

As presented in Table 1, H\(_2\)O\(_2\) exposure reduced MMP, increased DNA fragmentation and Ca\(^{2+}\) release in NGF differentiated-PC12 cells (P < 0.05). GAE pre-treatments reversed these changes (P < 0.05), in which concentration-dependent effects were presented in increasing MMP and reducing DNA fragmentation (P < 0.05). H\(_2\)O\(_2\) exposure enhanced caspase-3 and caspase-8 activities; and lowered Na\(^+\)-K\(^+\)-ATPase activity in NGF differentiated-PC12 cells (Fig. 4, P < 0.05). GAE pre-treatments at test concentrations decreased caspase-3 activity and increased Na\(^+\)-K\(^+\)-ATPase activity (P < 0.05). However, GAE pre-treatment only at 1% reduced caspase-8 activity (P < 0.05).

#### 3.3. GAE mitigated oxidative and inflammatory stress

As presented in Table 2, H\(_2\)O\(_2\) stimulated ROS generation, decreased GSH content, and reduced GPX and catalase activities in NGF differentiated-PC12 cells (P < 0.05). GAE pre-treatments reversed these changes (P < 0.05), in which concentration-dependent effects were presented in increasing MMP and reducing DNA fragmentation (P < 0.05). H\(_2\)O\(_2\) exposure enhanced caspase-3 and caspase-8 activities; and lowered Na\(^+\)-K\(^+\)-ATPase activity in NGF differentiated-PC12 cells (P < 0.05). GAE pre-treatments at test concentrations decreased caspase-3 activity and increased Na\(^+\)-K\(^+\)-ATPase activity (P < 0.05). However, GAE pre-treatment only at 1% reduced caspase-8 activity (P < 0.05).
Table 1 – Effects of GAE upon MMP, measured as MFI; DNA fragmentation, measured as enrichment factor; and Ca\(^{2+}\) release. NGF differentiated-PC12 cells were pre-treated with GAE at 0.25%, 0.5% or 1%, and followed by using H\(_2\)O\(_2\) to induce cell injury. Normal group had no GAE or H\(_2\)O\(_2\). Control group had no GAE, but with H\(_2\)O\(_2\). Data are mean ± SD (n = 7). *Values in a column without a common letter differ, P < 0.05.

|                  | MFI      | enrichment factor | [Ca\(^{2+}\)], nM |
|------------------|----------|-------------------|-------------------|
| Normal           | 100\(^{a}\) | 1.00\(^{a}\)      | 461 ± 49\(^{a}\)  |
| Control          | 28 ± 2\(^{a}\) | 2.31 ± 0.12\(^{a}\) | 1618 ± 143\(^{a}\) |
| GAE, 0.25        | 39 ± 4\(^{a}\) | 2.01 ± 0.09\(^{a}\) | 1290 ± 98\(^{a}\)  |
| GAE, 0.5         | 54 ± 6\(^{a}\) | 1.69 ± 0.10\(^{a}\) | 1007 ± 56\(^{a}\)  |
| GAE, 1           | 68 ± 3\(^{a}\) | 1.4 ± 0.07\(^{a}\)  | 927 ± 48\(^{a}\)   |

Fig. 1 - Effects of GAE upon cell viability (a) and plasma membrane damage (b) without H\(_2\)O\(_2\) treatment. NGF differentiated-PC12 cells were treated with GAE at 0.25%, 0.5% or 1%. Normal group had no GAE. Data are mean ± SD (n = 7). *Values among bars without a common letter differ, P < 0.05.

Fig. 2 - Effects of GAE upon cell viability (a) and plasma membrane damage (b) with H\(_2\)O\(_2\) treatment. NGF differentiated-PC12 cells were pre-treated with GAE at 0.25%, 0.5% or 1%, and followed by using H\(_2\)O\(_2\) to induce cell apoptosis. Normal group had no GAE or H\(_2\)O\(_2\). Control group had no GAE, but with H\(_2\)O\(_2\). Data are mean ± SD (n = 7). **Values among bars without a common letter differ, P < 0.05.
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4. Discussion

The data of our present work revealed that without \( \text{H}_2\text{O}_2 \) stimulation, GAE treatments at test concentrations did not affect viability and plasma membrane integrity of NGF differentiated-PC12 cells. These findings implied that GAE might not have adverse impact for neuronal cells. Our previous animal study reported that dietary GAE intake at 0.5% markedly attenuated ethanol-induced hepatic glycative damage and lipid accumulation [13]. Our current cell line study found that GAE at three test concentrations did not affect viability, which in turn enhanced anti-apoptotic defense and improved cell viability. These results suggest that GAE could mediate Bcl-2/Bax pathway and increase cell survival. On the other hand, \( \text{H}_2\text{O}_2 \) exposure impaired plasma membrane integrity and caused DNA fragmentation, which definitely contributed to cell rupture and apoptosis [18, 19]. Our data revealed that GAE reversed these alterations. These findings indicated that GAE could benefit DNA stability and maintain the integrity of plasma membranes, which consequently favored cell survival. In addition, GAE pre-treatments diminished \( \text{H}_2\text{O}_2 \) induced Ca\(^{2+}\) release observed in NGF differentiated-PC12 cells. The less Ca\(^{2+}\) release observed in GAE treated NGF differentiated-PC12 cells could be partially ascribed to the improvement from GAE upon plasma membrane integrity. It is reported that released Ca\(^{2+}\) facilitates nerve impulse transmission and stimulates neuronal excitability, which might promote the development and progression of seizure [20, 21]. Our data implied that GAE might decrease neuronal excitability through limiting Ca\(^{2+}\) release. Further study regarding the anti-seizure effect of GAE is worthy to be investigated.

Table 2 – Effects of GAE upon ROS and GSH levels, and GPX and catalase activities. NGF differentiated-PC12 cells were pre-treated with GAE at 0.25%, 0.5% or 1%, and followed by using \( \text{H}_2\text{O}_2 \) to induce cell injury. Normal group had no GAE or \( \text{H}_2\text{O}_2 \). Control group had no GAE, but with \( \text{H}_2\text{O}_2 \). Data are mean ± SD (n = 7). **Values in a column without a common letter differ, \( P < 0.05 \).

|                | ROS RFU/mg protein | GSH ng/mg protein | GPX U/mg protein | catalase U/mg protein |
|----------------|--------------------|-------------------|------------------|----------------------|
| Normal         | 0.14 ± 0.06\(^a\)  | 90 ± 5\(^c\)      | 66 ± 4\(^a\)     | 2.52 ± 0.19\(^a\)    |
| Control        | 2.36 ± 0.13\(^e\)  | 39 ± 4\(^e\)      | 38 ± 2\(^e\)     | 0.97 ± 0.08\(^e\)    |
| GAE, 0.25      | 2.00 ± 0.05\(^d\)  | 49 ± 3\(^b\)      | 45 ± 3\(^b\)     | 1.27 ± 0.11\(^b\)    |
| GAE, 0.5       | 1.58 ± 0.1\(^c\)   | 62 ± 2\(^c\)      | 52 ± 4\(^c\)     | 1.58 ± 0.12\(^c\)    |
| GAE, 1         | 1.17 ± 0.09\(^d\)  | 66 ± 3\(^c\)      | 55 ± 3\(^d\)     | 1.89 ± 0.07\(^d\)    |

mRNA expression (\( P < 0.05 \)).

Fig. 3 - Effects of GAE upon mRNA expression of Bcl-2 and Bax. NGF differentiated-PC12 cells were pre-treated with GAE at 0.25%, 0.5% or 1%, and followed by using \( \text{H}_2\text{O}_2 \) to induce cell apoptosis. Normal group had no GAE or \( \text{H}_2\text{O}_2 \). Control group had no GAE, but with \( \text{H}_2\text{O}_2 \). Data are mean ± SD (n = 7). **Values among bars without a common letter differ, \( P < 0.05 \).

Fig. 4 - Effects of GAE upon the activity of caspase-3, caspase-8 and Na\(^+\)-K\(^+\)-ATPase. NGF differentiated-PC12 cells were treated with GAE at 0.25%, 0.5% or 1%, and followed by using \( \text{H}_2\text{O}_2 \) to induce cell injury. Normal group had no GAE or \( \text{H}_2\text{O}_2 \). Control group had no GAE, but with \( \text{H}_2\text{O}_2 \). Data are mean ± SD (n = 7). **Values among bars without a common letter differ, \( P < 0.05 \).
Collapse of MMP activates apoptotic executors such as caspase-3 and caspase-8 [22]. The raised activity of these two caspases further induced alterations in cellular morphological characteristics and nuclear protein cleavage, and all these events led to cell death [23]. Na\(^{+}\)-K\(^{+}\)-ATPase, a transmembrane protein, is in charge of intracellular Na\(^{+}\) exchange for extracellular K\(^{+}\). The loss of MMP caused the reduction in Na\(^{+}\)-K\(^{+}\)-ATPase activity, which subsequently impaired ion homeostasis and promoted apoptotic insult [24]. Unterberg et al. [25] indicated that lower Na\(^{+}\)-K\(^{+}\)-ATPase activity contributed to neuronal swelling and even brain edema. In our present study, H\(_2\)O\(_2\) exposure disturbed mitochondrial membrane, which was evidenced by greater caspase-3 and caspase-8 activities, as well as lower Na\(^{+}\)-K\(^{+}\)-ATPase activity. However, GAE pre-treatments attenuated mitochondrial membrane injury caused by H\(_2\)O\(_2\). One possibility was that GAE enhanced the defensive capability of mitochondrial membrane against H\(_2\)O\(_2\), which consequently diminished the impact from H\(_2\)O\(_2\) upon caspase-3, caspase-8 and Na\(^{+}\)-K\(^{+}\)-ATPase activity. The other possibility was that GAE directly affected caspases and Na\(^{+}\)-K\(^{+}\)-ATPase activities, which finally mitigated apoptotic stress and benefited Na\(^{+}\)/K\(^{+}\) ion homeostasis. These data once again suggest that GAE could maintain mitochondrial membrane stability and alleviated apoptotic stress in H\(_2\)O\(_2\)-treated NGF differentiated-PC12 cells.

It is reported that GAE contained many phytochemicals with anti-oxidative and anti-inflammatory activities such as ferulic acid, chlorogenic acid, quercetin and apigenin [12]. Actually, the anti-oxidative and anti-inflammatory protection of ferulic acid and quercetin for neuronal cells or brain tissue has been reported [26, 27]. Thus, the less production of ROS and inflammatory cytokines, greater GSH content, greater GPX and catalase activities as we observed in GAE treated NGF differentiated-PC12 cells could be ascribed to the presence of phytochemicals in GAE. We believe that the mitigated oxidative and inflammatory stress also contributed to stabilize DNA and mitochondrial membrane integrity in GAE treated NGF differentiated-PC12 cells, which in turn improved cell survival. As observed by others, the activation of NF-xB and p38 signaling pathways due to H\(_2\)O\(_2\) stimulation facilitated the generation of oxidants and inflammatory factors such as ROS and TNF-alpha [28, 29]. Our data agreed that H\(_2\)O\(_2\) was a promotor responsible for neuronal cell oxidative and inflammatory injury. However, our findings indicated that GAE pre-treatments limited the mRNA expression of NF-xB and p38. It is highly possible that the pre-treatments of GAE led to some phytochemical components penetrate plasma membrane of NGF differentiated-PC12 cells, where these component compounds exert their protective actions against subsequent H\(_2\)O\(_2\) assault. Since these signaling pathways have been suppressed, the lower production of downstream factors such as ROS and inflammatory cytokines could be explained. These finding also suggest that GAE was able to protect NGF differentiated-PC12 cells at molecular levels.

*G. bicolor* is a vegetable. Its aqueous extract is easily prepared and should be safe. Moreover, our previous animal study reported that dietary GAE protected liver against ethanol induced injury [13]. This animal study supported that the active component compounds of GAE could be absorbed, metabolized and exerted its bioactivities. However, it remains unknown that GAE could pass blood brain barrier, and protect brain or neurons. Further animal study is definitely necessary to verify the protective effects of GAE upon brain or neurons. In addition, the phytochemical profile of GAE might not be consistent due to environmental factors such as seasons and planting conditions. Thus, standardization process is very important for the used GAE.

In conclusion, aqueous extract of G. bicolor leaf part enhanced NGF differentiated-PC12 cells survival against H\(_2\)O\(_2\) through maintaining mitochondrial membrane potential, decreasing oxidative and inflammatory injury, and regulating the mRNA expression of Bcl-2, Bax, NF-xB and p38. These findings sug-

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**Table 3 — Effects of GAE upon level (pg/mg protein) of IL-1beta, IL-6 and TNF-alpha.** NGF differentiated-PC12 cells were pre-treated with GAE at 0.25%, 0.5% or 1%, and followed by using H\(_2\)O\(_2\) to induce cell injury. Normal group had no GAE or H\(_2\)O\(_2\). Control group had no GAE, but with H\(_2\)O\(_2\). Data are mean ± SD (n = 7). *a-dValues among bars without a common letter differ, P < 0.05.

| GAE (mg/mL) | IL-1beta | IL-6 | TNF-alpha |
|------------|----------|------|-----------|
| Normal     | 10 ± 5\(^a\) | 9 ± 3\(^b\) | 8 ± 4\(^d\) |
| Control    | 76 ± 7\(^d\) | 79 ± 5\(^d\) | 91 ± 6\(^d\) |
| GAE, 0.25  | 62 ± 4\(^c\) | 60 ± 6\(^a\) | 75 ± 4\(^b\) |
| GAE, 0.5   | 48 ± 2\(^b\) | 46 ± 4\(^b\) | 58 ± 3\(^d\) |
| GAE, 1     | 45 ± 4\(^b\) | 41 ± 3\(^b\) | 42 ± 4\(^d\) |

**Fig. 5 - Effects of GAE upon mRNA expression of NF-kB and p38.** NGF differentiated-PC12 cells were treated with GAE at 0.25%, 0.5% or 1%, and followed by using H\(_2\)O\(_2\) to induce cell injury. Normal group had no GAE or H\(_2\)O\(_2\). Control group had no GAE, but with H\(_2\)O\(_2\). Data are mean ± SD (n = 7). a-dValues among bars without a common letter differ, P < 0.05.
gested that this aqueous extract might possess neuronal protective potential.

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Conflict of interest statement

The authors wish to disclose no conflicts of interest.

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