Introduction
Alzheimer’s disease (AD) and Parkinson’s disease (PD) are multifaceted, progressive neurodegenerative disorders that occur mainly in older age groups; however, the incidence of these diseases has increased in younger populations in recent years. Although these two disorders are characterized by different clinical manifestations and pathological phenomena, they are both categorized by a selective loss of neurons. Free radical-induced oxidative stress, mitochondrial dysfunction, and excitotoxic processes have been suggested to play a key role in the loss of neurons[1, 2].

Hsp70, a member of the heat shock protein family, functions as an ATP-dependent molecular chaperone in protein folding, multi-protein complex assembly, transmembrane transport, and protein degradation[3]. This protein is also part of an inducible system that aids in cell survival by preventing and/or repairing stress-induced protein damage during and after detrimental environmental stresses, such as free-radical accumulation[4]. Pytlowany et al used PC12 cells to compare the protective effects of huperzine A (HupA) derivative M3 and HupA on sodium nitroprusside (SNP)-induced apoptosis[5]. PC12 cells are derived from a rat pheochromocytoma cell line and can acquire neuron-like properties, such as neurite extension, when exposed to nerve growth factor (NGF). These cells are often used to study PD because NGF-treated and untreated cells synthesize, store, secrete, and take up dopamine by processes that are similar to those of dopaminergic neurons[6]. PC12 cells are also used to study AD because they express β-amyloid (Aβ) precursor protein (APP), in keeping with the mRNA and protein levels found in cortical neurons[7].

HupA, which is isolated from the herb *Huperzia serrata*, has been shown to be a highly effective, selective, and reversible acetylcholinesterase (AChE) inhibitor. The use of HupA as a therapeutic treatment for early or mild stages of AD may be due to additional mechanisms other than the reversible inhibition of AChE in the central nervous system. For example, HupA may target oxidative stress, Aβ-associated neurotoxicity, APP processing, and NGF[8]. The capacity of AChE to hydrolyze acetylcholine is not involved in the aforementioned mechanisms[9]. Therefore, we sought to identify derivatives of HupA that retain a similar level of AChE inhibition but yield an increase in cell protection. Due to the rigid molecular
configuration of HupA, traditional chemical methods cannot be used to modify its structure. Instead, we chose to use microbial transformation to produce HupA derivatives. M3 is a product of the transformation of HupA by Streptomyces griseus CACC 200300 after a two-step procedure and various chromatographic techniques. The structure (Figure 1) was identified as huperzine A 8α,15α-epoxide using physical and chemical data collected through multiple analyses, such as HRMS, 1D NMR, 2D NMR, and IR. Although the IC₅₀ value of M3 on AChE inhibition was established to be higher than that of HupA[10], the remarkable effect of protecting PC12 cells from oxidative stress was observed in our present screening.

Figure 1. The structures of HupA and M3 by Streptomyces griseus CACC 200300.

Materials and methods

Materials and reagents

M3 was prepared as previously described[10]. HupA and SNP were bought from the National Institute of the Control of Pharmaceutical and Biological Products. Quercetin and Hoechst 33342 were purchased from Sigma. These compounds were dissolved in DMSO and diluted by PC12 medium. An annexin V/propidium iodide (PI) staining kit was bought from Beijing Biosea Biotechnology Company. Dichlorofluorescein diacetate (DCFH-DA) and MDA kits were bought from Beyotime Institute of Biotechnology. The anti-Hsp70 antibody and anti-mouse secondary antibody were bought from Santa Cruz Biotechnology. The anti-caspase-3 (17 kDa) antibody was bought from Cell Signaling Technology. The anti-β-actin antibody was bought from Sigma.

Cell culture and treatment

PC12 cells, purchased from the American Tissue Type Collection, were grown in 75-cm² tissue culture flasks in DMEM supplemented with 5% heat-inactivated fetal bovine serum (FBS), 10% equine serum (ES), penicillin (100 IU/mL), streptomycin (100 mg/L), and L-glutamine (2 mmol/L). Cultures were maintained at 37°C in 95% air-5% CO₂ in a humidified incubator and passaged every 3 to 4 d. PC12 cells in logarithmic phase were seeded into 96-well or 6-well plates coated with 0.1 mg/mL poly-L-lysine (Sigma) and allowed to grow for at least 24 h. Different concentrations of M3 and HupA (0.1, 1, and 10 μmol/L) were added to the culture media for 1 h followed by addition of SNP (200 μmol/L) for 24 h. In some experiments, PC12 cells were pretreated with 10 μmol/L quercetin and 10 μmol/L M3 for 1 h and were exposed to 200 μmol/L SNP for the next 24 h.

MTT assay

PC12 cells were cultured at a density of 6000 cells per well in 96-well plates. MTT (5 mg/mL) was added to each well after the aforementioned treatments and the plates were incubated for 4 h. After centrifugation, the supernatant was removed from each well, and DMSO (100 μL) was added. After 30 min, the absorbance was recorded with a microplate reader (Bio-Rad model 550) at a wavelength of 570 nm. Each experiment was performed in triplicate.

Detection of apoptosis

PC12 cells were cultured at a density of 30000 cells per well in 6-well plates. Cells were harvested by being digested in 0.25% trypsin after treatment, stained with annexin V for 20 min, and PI for another 10 min at 37°C before detection using flow cytometry (EPICS XL, Beckman Coulter, USA). To stain the DNA, cells were incubated with Hoechst 33342 (10 μg/mL) diluted in phosphate-buffered saline (PBS) for 10 min at room temperature. Cells were then observed by the IN Cell Analyzer 1000 (GE Healthcare, USA).

Measurement of reactive oxygen species (ROS)

Intracellular accumulation of ROS was measured using fluorophotometric quantitation. The cells were cultured with M3 or HupA (10 μmol/L) for 1 h before exposure to SNP (200 μmol/L) for 24 h. The harvested cells were subsequently stained with DCFH-DA for 30 min at 37°C. The cell suspension was dispensed into 96-well black plates. DCFH-DA reacts with ROS and is converted to dichlorofluorescein (DCF). The fluorescence from the DCF was analyzed using a Spectra microplate fluorescence reader (Max Gemini XS, MD, USA) with the excitation wavelength set at 488 nm and the emission wavelength set at 525 nm.

Estimation of lipid peroxidation

MDA reacts with thiobarbituric acid (TBA) to produce a florescent product. Levels of MDA were measured in PC12 cell lysates with a microplate reader at a wavelength of 535 nm. The cells were cultured with M3 or HupA (10 μmol/L) for 24 h. After centrifugation, the supernatant was removed, and DMSO (100 μL) was added. After 30 min, the absorbance was recorded with a microplate reader (Bio-Rad model 550) at 570 nm. Each experiment was performed in triplicate.

Protein assay

After a 1-h treatment with M3 and 24 h exposure to SNP, cells were rapidly washed with PBS and lysed on ice in a lysis buffer (pH=8.0) containing 50 mmol/L Tris·HCl, 150 mmol/L NaCl, 1% NP-40, 1 mmol/L PMSF, 1 mmol/L EDTA, 50 mmol/L NaF, 1 mmol/L DTT, 1 mg/L aprotinin and 1% Triton X-100. Protein concentrations were measured with a BCA kit (Vigorous). The lysates were solubilized in an SDS sample buffer, separated with 15% SDS-PAGE, and transferred
to PVDF membrane (Millipore). The membrane was blocked with 3% BSA and subsequently incubated with anti-Hsp70, anti-caspase-3 (17 kDa), or anti-β-actin antibodies followed by a horseradish peroxidase (HRP)-conjugated secondary antibody and detected with an ECL detection system (Molecular Device, Lmax). The density of each band was quantitated using image-analysis software (Science Lab 2005 Image Gauge; Fuji Film Co Ltd, Tokyo, Japan).

Statistical analysis
Data are expressed as the mean±standard deviation. An analysis of variance (ANOVA) followed by a Newman-Keuls post-hoc test was performed to assess the differences between groups. Values of $P<0.05$ were regarded as statistically significant.

Results
The neurotoxicity of SNP on PC12 cells
Before studying the protective effects of M3, we analyzed the toxicity of SNP on PC12 cells. PC12 cells treated with different concentrations of SNP for 24 h displayed a decrease in the production of formazan dye in a dose-dependent manner. Exposure to 200 μmol/L of SNP induced moderate injury to the cells when compared with the control group. This concentration of SNP was used to compare the neuroprotective effects of M3 and HupA (Figure 2).

Pre-incubation with M3 and HupA protected PC12 cells from SNP-induced injury
Exposure to 200 μmol/L of SNP for 24 h decreased the viability of PC12 cells to 69.0%±4.8% compared with the control group. Pre-incubation with M3 or HupA for 1 h attenuated the injury induced by SNP in a dose-dependent manner. This effect was statistically significant at a concentration of 10 μmol/L ($P<0.05$). The cell viability after pre-incubation with M3 was 84.1%±2.7% of the control, while the cell viability after pre-incubation with HupA was 77.1%±3.5% of the control ($P<0.05$, Figure 3). There was no significant difference in cell viability between the M3 and HupA pre-treated groups ($P>0.05$).

M3 and HupA attenuated the SNP-induced apoptosis of PC12 cells
Analysis of an annexin V/PI stain using flow cytometry and Hoechst 33342 stain with a fluorescence microscope revealed that pretreatment with HupA or M3 significantly decreased the apoptosis of PC12 cells induced by SNP. M3 pretreatment alone showed no effect on apoptotic bodies or the nuclear morphology of PC12 cells. Pre-treatment with HupA and M3 (10 μmol/L) attenuated nuclear fragmentation and chromatin condensation, and inhibited apoptosis at the early and intermediate stages of SNP exposure. There was no statistically significant difference in the percentage of apoptotic bodies between the M3 and HupA groups (Figure 4).

M3 inhibited the production of SNP-induced ROS in PC12 cells
DCFH-DA is hydrolyzed to DCFH by intracellular esterases, which is subsequently oxidized by ROS to DCF, which is a fluorescent compound. The fluorescence intensity is directly proportional to the amount of ROS generated from SNP exposure. Using an excitation wavelength of 488 nm and an emission wavelength of 525 nm, the mean fluorescence intensity increased to 183.5%±18.3% in the SNP-treated group compared with the control group. Pre-incubation with 10 μmol/L of M3 or 10 μmol/L of HupA resulted in a decrease in the mean fluorescence intensity increased to 183.5%±18.3% in the SNP-treated group compared with the control group. Pre-incubation with 10 μmol/L of M3 or 10 μmol/L of HupA resulted in a decrease in the mean fluorescence intensity to 142.3%±20.2% or 132.0%±6.5%, respectively, compared with the SNP-treated group (Figure 5). There was no statistically significant difference in the levels of ROS between the M3 and HupA groups ($P>0.05$).

Treatment with M3 decreased SNP-induced MDA accumulation in PC12 cells
MDA, formed by degradation of polyunsaturated lipids by

Figure 2. Effect of SNP on the cell viability in PC12 cells. Cells were exposed to SNP for 24 h. The cell viability was evaluated by measuring the quantity of formazan generated by activated mitochondria. The values were expressed as percentage of control, which is set to 100%. The percentage of cell viability was presented as mean±SD for six replicates. $b P<0.05$ vs control group.

Figure 3. Effect of HupA and M3 on the cell viability in PC12 cells induced by SNP. PC12 cells were pre-treated with HupA and M3 at different concentrations for 1 h before 200 μmol/L SNP was added. The percentage of cell viability was presented as mean±SD for six replicates. $b P<0.05$ vs control group. $e P<0.05$ vs SNP group.
ROS, is used as a marker to measure the level of oxidative stress in an organism. In our previous study, SNP increased MDA levels to 382.4%±57.2% compared with the control group. However, pre-incubation with 10 μmol/L of M3 or 10 μmol/L of HupA for 1 h caused a decrease in the levels of MDA to 288.7%±21.6% in the HupA pretreated group and 246.9%±44.2% in the M3 pretreated group (Figure 5). There was no statistically significant difference between the level of MDA in the M3 and HupA groups (P>0.05).

**Treatment with M3 increased the expression of Hsp70 in PC12 cells**

It was reported that an increase in the expression of Hsp70 confers an improvement in the survival ability of cells against SNP toxicity. The effect of M3 on the expression of Hsp70 in PC12 cells was analyzed in this study. The expression of Hsp70 increased in a time-dependent manner after M3 exposure, peaking after 0.5 h. No change was observed in Hsp70 expression after pre-incubation with HupA. After 1 h of exposure to SNP, the expression of Hsp70 increased. Pretreatment with M3 increased the Hsp70 level compared with cells exposed to SNP (Figure 6).

**Quercetin blocked the protective effect of M3 treatment on SNP-induced apoptosis in PC12 cells**

Quercetin, a HSP biosynthesis inhibitor, was used to confirm if Hsp70 was involved in the protective effect of M3. As shown in Figure 7A, quercetin attenuated the protection conferred from M3 pretreatment on SNP-induced apoptosis while showing almost no effect on HupA-treated cells. As displayed in
Figure 7B, treatment with HupA failed to significantly change the expression of Hsp70 compared with the expression of the SNP-exposed group. In M3 pretreated cells, the expression of Hsp70 increased to 212.5%±26.7% compared with control group. Caspase-3 expression remarkably increased in PC12 cells after exposure to SNP for 24 h, and pre-incubation with HupA and M3 for 1 h reversed this effect. Quercetin had no effect on the activation of caspase-3 in the control or HupA pretreated group, but it significantly increased the expression of caspase-3 in the M3 pretreated group (P<0.05).

Discussion

Oxidative stress is recognized as an important mechanism in several neurodegenerative diseases. It has been proven that the activity of inducible nitric oxide synthases (iNOS) was increased in the brains of AD patients\cite{11}. Free-radical production and nitration of endogenous biomolecules are the two main reasons that accumulation of nitric oxide (NO) induces cell loss\cite{12}. In this study, SNP was used to induce the apoptosis of PC12 cells in order to evaluate potential neuroprotective compounds. Exposure to SNP in vitro could partially mimic the oxidative stress observed in the brains of neurodegenerative patients\cite{5}. We found that, similar to the effect of HupA, pretreatment with M3 attenuated the apoptosis induced by SNP in PC12 cells. Although M3 and HupA showed a similar level of ROS scavenging, it seems that this was not the only mechanism involved in the protective effects of M3. Indeed, treatment with M3 induced the expression of Hsp70 in a time-dependent manner, unlike HupA. Hsp70, a member of the...
Overproduction of Hsp70 was shown to strongly protect brain cells from ischemia/reperfusion injuries both in vitro and in vivo[13]. Some of the anti-apoptotic activity of Hsp70 could be attributed to its ability to suppress the activity of JUN-kinase or Bid proteins[16, 17]. Thus, the decreased activation of caspase-3 in the M3 treated group might be due to the inhibition of the JNK/Bim pathway by the induction of Hsp70.

NO exposure has been established to induce Hsp70 expression in order to protect cells from tumor necrosis factor-induced apoptosis[18]. In our study, expression of Hsp70 could not save the PC12 cells from SNP-induced apoptosis. However, the M3 pretreatment remarkably increased the Hsp70 response without causing detrimental effects.

To confirm whether the induction of Hsp70 by M3 was responsible for its protective effects, quercetin, which is an HSP biosynthesis inhibitor, was used. Our results showed that pretreatment with quercetin attenuated the protective effect of M3 against SNP-induced apoptosis. This finding suggests that the protective effect of M3 was mediated, at least partly, by increasing the expression of Hsp70. However, quercetin failed to attenuate the protection of HupA on SNP-induced apoptosis, suggesting that the protective mechanism of HupA might have little interaction with Hsp70. Therefore, it appears that the protective mechanisms for these two compounds are different.

Taken together, M3 was similar to HupA with respect to free-radical scavenging. However, M3 treatment was able to elevate the expression of Hsp70 and decrease the expression of active caspase-3. M3 may activate the stress response without causing detrimental effects, thereby allowing the cells to survive under lethal conditions. Pretreatment with HupA also displayed ROS scavenging in PC12 cells exposed to SNP, but the presence of other protective mechanisms is not clear. Our research suggests that the distinctive chemical structure of HupA and its derivatives may be related to its anti-apoptotic properties. Further studies will concentrate on optimizing and analyzing these chemical structures in order to facilitate the design of novel anti-neurodegenerative drugs.

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Author contribution
Na NING, Jin-feng HU, Yu-he YUAN, and Nai-hong CHEN developed the research plan; Na NING performed the experiments; Xin-yuan ZHANG and Jun-gui DAI contributed reagents or analytical tools; Na NING and Nai-hong CHEN analyzed the data; and Na NING prepared the manuscript.

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Figure 7. Quercetin inhibited the protective effect of M3 against SNP-induced apoptosis in PC12 cells. PC12 cells were pre-treated with 10 μmol/L HupA, M3, or 10 μmol/L quercetin for 1 h then exposed to 200 μmol/L SNP for the next 24 h. (A) Effect of quercetin on the cell viability in PC12 cells by MTT method. The percentage of cell viability was presented as mean±standard deviation for six replicates. (B) Effect of quercetin on the ratio of Caspase-3 and Hsp70 in PC12 cells. The percentage of density was presented as mean±standard deviation for three individual experiments. *P<0.01 vs control group. †P<0.05 vs SNP group. ‡P<0.05 vs M3 pretreated group.

heat shock protein family, functions as a chaperone in protein folding, which can reduce abnormal protein aggregation[13, 14].

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