Mulberry fruit protects dopaminergic neurons in toxin-induced Parkinson’s disease models

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Parkinson’s disease (PD), one of the most common neurodegenerative disorders, is characterised by the loss of dopaminergic neurons in the substantia nigra pars compacta (SNpc) to the striatum (ST), and involves oxidative stress. Mulberry fruit from Morus alba L. (Moraceae) is commonly eaten, and has long been used in traditional oriental medicine. It contains well-known antioxidant agents such as anthocyanins. The present study examined the protective effects of 70% ethanol extract of mulberry fruit (ME) against neurotoxicity in in vitro and in vivo PD models. In SH-SY5Y cells stressed with 6-hydroxydopamine (6-OHDA), ME significantly protected the cells from neurotoxicity in a dose-dependent manner. Other assays demonstrated that the protective effect of ME was mediated by its antioxidant and anti-apoptotic effects, regulating reactive oxygen species and NO generation, Bcl-2 and Bax proteins, mitochondrial membrane depolarisation and caspase-3 activation. In mesencephalic primary cells stressed with 6-OHDA or 1-methyl-4-phenylpyridinium (MPP⁺), pre-treatment with ME also protected dopamine neurons, showing a wide range of effective concentrations in MPP⁺-induced toxicity. In the sub-acute mouse PD model induced by 1-methyl-4-phenyl-1,2,3,6-tetrahydrodipyridine (MPTP), ME showed a preventative effect against PD-like symptoms (bradykinesia) in the behavioural test and prevented MPTP-induced dopaminergic neuronal damage in an immunocytochemical analysis of the SNpc and ST. These results indicate that ME has neuroprotective effects in in vitro and in vivo PD models, and that it may be useful in preventing or treating PD.

Mulberry fruit: Parkinson’s disease: Neuroprotective effects: 6-Hydroxydopamine: 1-Methyl-4-phenyl-1,2,3,6-tetrahydrodipyridine

Worldwide, as the population ages, the probability of people developing neurodegenerative diseases such as Alzheimer’s disease and Parkinson’s disease (PD) is increasing. PD is the second most common neurological disorder, affecting 2% of the population over the age of 60 years; it is characterised by the selective loss of dopaminergic neurons in the substantia nigra pars compacta (SNpc)(1). Although the cause of PD remains unclear, evidence from many studies suggests the involvement of oxidative stress(2,3). Oxidative stress is thought to be one of the factors that reduce cognitive and motor performance in neurodegenerative disease; oxidative defence mechanisms such as catalase and GSH decline, while oxidative damage molecules, such as hydroxyl radicals and peroxynitrite, increase. Thus, a high level of antioxidant activity has been linked with protection against neurodegenerative diseases(23).

Various berries have long been reported to have preventative or positive effects in many disease states including cancer(4), CVD(5) and age-related brain diseases(6). Berries contain numerous phytochemicals such as flavonoids, tannins and phenolic acids, and recently, the antioxidant effects of berry components have been studied(7). Fruit-derived polyphenolic compounds are known to have antioxidant properties and to potentially be neuroprotectants(8–10). Moreover, research has revealed that some berries have neuroprotective effects; strawberries and blueberries were shown to reverse declines in cognition, motor behaviour, neuronal signal transduction and memory; furthermore, blueberries showed neuroprotective effects in an Alzheimer’s disease model(11–16). Shukitt-Hale et al. (17) reported that berry fruits may lower the risk of developing age-related neurodegenerative diseases, suggesting that...
polyphenolic compounds in berry fruits may reduce oxidative stress and inflammation.

Mulberry (Morus alba L.), a member of the Moraceae family, has been naturalised and widely cultivated; mulberry fruit is commonly eaten, often dried, or made into wine. In traditional oriental medicine, it has been used to treat premature grey hair, to nourish the blood, to treat constipation and diabetes and to generate body fluids, which generally means enhancing health and promoting longevity. In addition, it has been reported to ameliorate inflammation-related haematological parameters in carrageenan-induced arthritic rats, to promote recovery from physical stress, to have neuroprotective effects against cerebral ischaemia and to have radical-scavenging properties.

Like other berry fruits, mulberry fruit contains not only high amounts of anthocyanins, a subset of the flavonoids that are important natural antioxidants, but also non-anthocyanin phenolics including rutin and quercetin known to have multi-bioactive functions including neuroprotective effects. It may be assumed that these compounds may have neuroprotective, antioxidant and anti-inflammatory effects in PD models. However, few studies have actually examined the possible neuroprotective outcomes in experimental settings. Thus, we examined the effects of the ethanol extract of mulberry fruit (ME) on dopaminergic neuron protection in in vitro PD models using the SH-SY5Y neuroblastoma with 6-hydroxydopamine (6-OHDA) and mesencephalic dopamine neurons stressed with 6-OHDA and 1-methyl-4-phenylpyridinium (MPP+). We also investigated the effects of ME in an in vivo PD model induced by 1-methyl-4-phenyl-1,2,3,6-tetrahydropyridine (MPTP).

**Experimental methods**

**Materials**

Dulbecco’s modified Eagle’s medium, minimal essential medium, fetal bovine serum and penicillin–streptomycin were purchased from Gibco Industries, Inc. (Auckland, New Zealand). 3-(4,5-Dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide (MTT), 6-OHDA, MPTP, MPP+, DMSO, Triton-X, glycine, JC-1 and 2,7-dichlorodihydrofluorescein (DCF) diacetate were purchased from Sigma-Aldrich (St Louis, MO, USA). Tetramethylrhodamine, protein assay reagent, Tween 20, ammonium persulphate, acrylamide, enzyme-linked chemiluminescence reagent and skimmed milk were purchased from Bio-Rad Laboratories (Hercules, CA, USA). Cyanidin-3-O-β-glucopyranoside was purchased from Polyspherol Laboratories AS (Sandnes, Norway). A caspase-3 assay kit and a mitochondria/cytosol fractionation kit were purchased from Assay Designs, Inc. (Ann Arbor, MI, USA). Biotinylated anti-rabbit antibody, normal goat serum and an avidin–biotin–peroxidase complex standard kit were purchased from Vector Laboratories (Burlingame, CA, USA).

**Preparation of extract**

Dried mulberry fruit was purchased from Jung Do Herbal Drug Company (Seoul, South Korea), and was extracted with 70% ethanol for 24 h at room temperature. Then, the extract was filtered, evaporated on a rotary vacuum evaporator and finally lyophilised. The powder (yield, 20.53%) was kept at 4°C. Before each experiment, the extract was dissolved in an appropriate vehicle and was vortex-mixed for 2 min at room temperature. ME was standardised based on the contents of C3g, one of the active components of mulberry, using reverse-phase HPLC (Agilent 1100 HPLC system; Agilent technologies, Inc., Santa Clara, CA, USA) equipped with a photodiode array detector. Separation was carried out using a J’sphere ODS-H80 column (250 × 4.6 mm, 4 μm; YMC Company Limited, Shin Kyoto, Japan) at 25°C. The mobile phases (A: 0.1% trifluoroacetic acid in acetonitrile and B: 0.1% trifluoroacetic acid in water) were 10–10% A for 0–10 min; 10–30% A for 10–20 min; and 30–100% A for 20–30 min at a flow rate of 1.0 ml/min. The detector wavelength was set at 520 nm. Four concentrations of C3g were prepared at 1, 2.5, 5 and 10 μg/ml, where 10 μl were injected as an external standard. ME and reference to the calibration curve obtained with C3g were analysed in triplicates. C3g was found in ME at a mean level of 0.43 (SEM 0.02) mg/g.

**Cell culture and treatment**

The SH-SY5Y cell line, a human neuroblastoma, was obtained from the Korea Cell Line Bank (Seoul, South Korea). Cells were maintained in Dulbecco’s modified Eagle’s medium, supplemented with 10% heat-inactivated fetal bovine serum, 100 units/ml penicillin and 100 μg/ml streptomycin in a water-saturated atmosphere of 5% CO2 at 37°C. The culture medium was changed every 3 d, and the cells were sub-cultured about twice a week. All experiments were carried out 48 h after the cells had been seeded in ninety-six-well plates and 60-mm dishes, at densities of 2 × 104 cells/ml and 2 × 106 cells/dish, respectively. After cells were about 80% confluent, various concentrations (0.1–100 μg/ml) of ME were added to the cells for 24 h at 37°C with or without 150 μM 6-OHDA for the last 6 h of ME treatment.

**Measurement of cell viability**

Treated cells were incubated with 1 mg/ml of MTT at 37°C in a CO2 incubator for 3 h. MTT medium was carefully aspirated from the wells, and the formazan dye was eluted using dimethyl sulphoxide. The plate was shaken on a shaker to dissolve the blue MTT-formazan. Absorbance was measured using a spectrophotometer (Versamax microplate reader, Molecular Device; Sunnyvale, CA, USA) at a wavelength of 570 nm, and then expressed as a percentage of the value in the vehicle-treated control culture.
Determination of intracellular reactive oxygen species

Intracellular ROS generation was measured using a fluorometer. DCFH diacetate passively enters the cells, and it is converted to non-fluorescent DCF. ROS react with DCFH to form the fluorescent product, DCF. The cells were seeded in ninety-six-well plates and were treated with ME and/or 6-OHDA. Then, the cells were incubated with 20 µM DCFH diacetate for 30 min. The fluorescence intensity was measured at 480 nm excitation and 530 nm emission using a fluorescence microplate reader (SpectraMax Gemini EM; Molecular Device).

Determination of extracellular nitric oxide

The accumulated level of nitrite (an indicator of NO) in culture supernatants was measured using the colorimetric reaction with the Griess reagent. The supernatants (100 µl) were transferred to a separate plate, and were reacted with 100 µl of Griess reagent in the dark for 10 min at room temperature. Absorbance at 550 nm was measured. For each experiment, freshly prepared NaNO₂ that had been serially diluted was used as a standard, in parallel with culture supernatants.

Assessment of mitochondrial membrane potential (Δψₘ)

The Δψₘ was measured using a fluorescent dye, JC-1 reagent. The treated cells were incubated with 1X JC-1 reagent solution at 37°C for 15 min. The red and green fluorescence were measured using a fluorescence microplate reader, with excitation at 585 and 510 nm and emission at 590 and 527 nm, respectively. The ratio of red to green fluorescence was quantified from the cells of interest.

Measurement of caspase-3 activation

The caspase-3 assay was performed according to the manufacturer’s protocol. Briefly, treated cell lysates were incubated with chilled lysis buffer on ice for 15 min. Then, they were centrifuged (14 000 g, 4 min, 4°C), and the supernatants were transferred to ninety-six-well plates. Reaction buffer containing 10 mM dithiothreitol was added to each well, and the 1 mM Asp-Glu-Val-Asp (DEVD) 7-amino-4-trifluoromethyl coumarin substrate was added and mixed. The mixture was incubated for 2 h at 37°C before protease activity was detected using a fluorescence microplate reader, with 360 nm excitation and 450 nm emission filters.

Western blot analysis

The treated cells were lysed with a triple-detergent lysis buffer to detect cleaved caspase-3. For the detection of Bax and Bcl-2, the treated cell lysates were incubated without (a) or with 150 µM 6-hydroxydopamine (6-OHDA) (b) for a further 6 h. Cell viabilities are expressed as a percentage of the controls (cells treated with vehicle for 24 h). Values are indicated as the mean values with their standard errors. *** Mean values were significantly different from the control group (P<0.001). ††† Mean values were significantly different from the ME-6-hydroxydopamine (6-OHDA)-induced accumulation of reactive oxygen species and nitric oxide in SH-SY5Y cells. The fluorescence intensity of the Griess reagent was measured after SH-SY5Y cells were exposed to 150 µM 6-OHDA for 1 h, followed by 20 µM DCFH diacetate for 30 min (a). Nitric oxide production in SH-SY5Y cells with ME pretreatment for 18 h before 150 µM 6-OHDA treatment was assayed by measuring the levels of nitrite in the supernatant fluid using the Griess reagent (b). Representative results from experiments. Values are indicated as the mean values with their standard errors. *** Mean values were significantly different from the control group (P<0.001). ††† Mean values were significantly different from the 6-OHDA-only treated group (P<0.001).
Bcl-2 proteins, the cells were fractionated into mitochondria and cytosol using a mitochondria/cytosol fractionation kit according to the manufacturer’s instructions. Cell lysates were separated by 15% SDS-PAGE, and were then transferred to a membrane. Membranes were incubated with 5% skimmed milk in Tris-buffered saline with Tween for 1 h and then

with the primary antibodies overnight at 4°C, followed by incubation with horseradish peroxidase-conjugated secondary antibodies for 1 h. Immunoreactive bands were detected using an ECL detection kit, and a LAS-4000 mini system (Fujifilm Corporation, Tokyo, Japan) was used for visualisation. The intensities of the bands were normalised to the β-actin band intensity using Multi Gauge software (Fujifilm Corporation).

**Cultures of mouse mesencephalic dopaminergic cells**

Cell cultures were prepared from the mesencephalon of 14 d embryos of timed pregnant Sprague–Dawley rats (Orient Bio, Osan, South Korea). Mesencephalons were dissected, collected, dissociated and plated in twenty-four-well plates with cover slips pre-coated with poly-L-lysine at a density of 1.5 × 10⁵ cells per well. Cultures were maintained in a humidified incubator of 5% CO₂ at 37°C in a minimal essential medium with 6·0 g/l glucose, 2 mM-glutamine and 10% fetal bovine serum. On day *in vitro* 6, the cells were treated with ME and were stressed with 10 µM-6-OHDA or 10 µM-MPP+ for a further 18 and 23 h, respectively. Then, the cells were fixed with 4% paraformaldehyde at room temperature for 30 min. The cells were stored in PBS at 4°C for immunocytochemistry.

**Animals**

Animal maintenance and treatment were carried out in accordance with the Principles of Laboratory Animal Care (NIH publication no. 85-23, revised 1985) and the Animal Care and Use guidelines of Kyung Hee University, Seoul, South Korea. Male C57Bl/6 mice (7 weeks) were purchased from Samtako, Inc. (Osan, South Korea). Animals were housed at an ambient temperature of 23 ± 1°C and at a relative humidity of 60 ± 10% under a 12 h light–dark cycle, and they were allowed free access to water and food.

**Drug administration**

Animals were assigned to three groups: (1) control group (*n* 8, intraperitoneally vehicle injected plus intraoarally vehicle-treated group); (2) MPTP group (*n* 8, intraperitoneally MPTP injected plus intraorally vehicle-treated group); (3) MPTP + ME group (*n* 8, intraperitoneally MPTP plus intraorally ME-treated group). ME that was dissolved in 10% dimethyl sulphoxide was administered at 500 mg/kg per d for 15 d. MPTP (MPTP base form) in normal saline was injected at 30 mg/kg per d for the last 5 d of ME treatment.

**Behavioural test and brain tissue preparation**

We performed the pole test, which measures motor coordination, on the seventh day after the last MPTP injection. The mice were placed head upward near the top of a vertical rough-surfaced pole (diameter 8 mm, height 55 cm). The time it took for the mice to turn completely downward (time to turn) and the time it took to reach the floor (locomotion activity time) were recorded, with a cut-off limit of 30 s. After the pole test, the mice were anaesthetised with 50 mg/kg Zoletil (intramuscularly) and were rapidly perfused transcardially with PBS, followed by 4% paraformaldehyde in 0·1 M-phosphate
buffer. Then, the brains were rapidly taken out, post-fixed in 4 % paraformaldehyde solution and processed for cryoprotection in 30 % sucrose at 4°C. Frozen brains were cut into 30-µm coronal sections using a cryostat microtome (CM3000; Leica, Wetzlar, Germany). Then, the tissues were stored in storing solution containing glycerine, ethylene glycol and phosphate buffer at 4°C for immunocytochemistry.

**Immunocytochemistry**

Mesencephalic cells on the cover slips and free-floating sections were rinsed in PBS at room temperature before immunostaining. They were pre-treated with 1 % H2O2 in PBS for 15 min to remove endogenous peroxidase activity. Then, they were incubated overnight at room temperature with a rabbit anti-TH antibody (1:2000 dilution) for dopaminergic neuron detection and a rabbit anti-Bax antibody (1:100 dilution) for apoptotic protein detection. They were then incubated with a biotinylated anti-rabbit IgG for 90 min, followed by incubation in avidin–biotin–peroxidase complex solution for 1 h at room temperature. The peroxidase activity was visualised with 3,3-diaminobenzidine for 3 min. After every incubation step, the cells and tissues were washed three times with PBS. Finally, the mesencephalic cells on the cover slips were mounted on gelatin-coated glass slides, air dried and photographed with a research microscope. The free-floating brain tissues were mounted on gelatin-coated slides, dehydrated, cleared with xylene and cover slipped using histomount medium. For quantification of the effect of ME in the mesencephalic dopaminergic cells, TH immunopositive cells were counted on at least four cover slips from independent experiments for each condition. Quantification of the effect of ME in brain tissues was performed by counting the TH-immunopositive cell number in SNpc at × 100 magnification under a microscope (AxioSkop 2; Carl Zeiss, Inc., Göttingen, Germany). The TH-immunopositivity in the striatum was measured at × 40 magnification using a StereoInvestigator (MBF Bioscience, Inc., Williston, ND, USA). Bax-immunopositive cells in SNpc were captured at × 200 magnification under a microscope. Data are presented as percentages of the control group values.

**Statistical analyses**

All quantitative data were analysed. The results are expressed as mean values with their standard errors. Statistical significance was determined by one-way ANOVA followed by the least significant difference test using SPSS (12.0K for Windows; Chicago, IL, USA). P values <0.05 were deemed to be statistically significant.

![Graphs and images](https://www.cambridge.org/core/core_media)
Results

Ethanol extract of mulberry fruit protects SH-SY5Y cells against 6-hydroxodopamine-induced neurotoxicity

To evaluate the protective activity of ME in SH-SY5Y cells, we used the MTT assay. Treatment with ME for 24 h at various concentrations had no influence on cell proliferation and caused no cell toxicity (Fig. 1(a)). Decreased cell viability induced by 150 µM 6-OHDA was reduced or prevented by ME pre-treatment at 0.1–100 µg/ml (Fig. 1(b)).

Ethanol extract of mulberry fruit inhibits 6-hydroxodopamine-induced stress in SH-SY5Y cells

To investigate whether the protective effect of ME on 6-OHDA-induced apoptosis involved ROS and NO generation, we used DCFH diacetate and the Griess reagent, respectively. Exposure to 150 µM 6-OHDA led to significant (P<0.001) ROS and NO elevation in SH-SY5Y cells, 151.86 (SEM 1.67)% and 468.76 (SEM 3.37)%, respectively, relative to the control. ME pre-treatment inhibited ROS (Fig. 2(a)) and NO (Fig. 2(b)) generation at 10 and 100 µg/ml and at 1, 10 and 100 µg/ml, respectively.

Ethanol extract of mulberry fruit inhibits 6-hydroxodopamine-induced apoptosis in SH-SY5Y cells

To determine the protective effects of ME on 6-OHDA-induced apoptosis, we measured Bax and Bcl-2 expression levels, ΔΨm and caspase-3 cleavage activity in SH-SY5Y cells. 6-OHDA-induced toxicity increased the Bax protein level in the mitochondria and decreased the Bcl-2 protein level, while ME protected SH-SY5Y cells from it (Fig. 3(a)). The ratio between green fluorescence (monomeric form, low ΔΨm) and red fluorescence (aggregated form, high ΔΨm) indicates the depolarisation of ΔΨm. 6-OHDA-induced toxicity significantly decreased ΔΨm (P<0.001), whereas ME pre-treatment at 10 and 100 µg/ml prevented depolarisation of the mitochondrial membrane (Fig. 3(b)). Caspase-3 activation was increased in the 6-OHDA group compared with the control group (P<0.001), whereas ME pre-treatment inhibited caspase-3 activity dose dependently (Fig. 3(c)); a maximal effect was obtained at 100 µg/ml ME. The expression level of cleaved caspase-3 in Western blot assay was consistent with its activity (Fig. 3(c)).

Ethanol extract of mulberry fruit has protective effects against 6-hydroxodopamine and 1-methyl-4-phenylpyridinium toxicity in mesencephalic dopaminergic neurons

To examine the protective effects of ME against 6-OHDA and MPP+ toxicity in primary dopaminergic neurons, we counted cell bodies with immunoreactivity to the anti-TH antibody. TH-positive cells were 500–700 cells per cover slip in control cultures. 6-OHDA neurotoxicity was defined as a 49-40 (SEM 1-15)% reduction in the survival rate, whereas pre-treatment with ME at 10 µg/ml protected dopaminergic cells, showing 65-04 (SEM 1-18)% (P<0.01) of TH-positive cells compared with the control group (Fig. 4(a)). MPP+ neurotoxicity was defined as a 48-04 (SEM 1-56)% reduction in the survival rate, whereas pre-treatment with ME also protected them, showing 69-55 (SEM 0-97)% (P<0.01) and 64-25 (SEM 4-50)% (P<0.05) of survival rates compared with the control group at 10 and 100 µg/ml, respectively (Fig. 4(b)).

Ethanol extract of mulberry fruit has neuroprotective effects against 1-methyl-4-phenyl-1,2,3,6-tetrahydropyridine toxicity in a mouse Parkinson’s disease model

We observed previously that ME protected mesencephalic dopaminergic neurons from 6-OHDA and MPP+ toxicity. In this experiment, we investigated whether ME could relieve motor symptoms and protect dopaminergic cells in the MPTP-induced mouse PD model. The pole test showed that the time to turn and locomotion activity time in the MPTP group were significantly prolonged compared with the control group, whereas they were significantly shortened in the MPTP+ME group (Fig. 5). Thus, ME prevented MPTP-induced bradykinesia at 500 mg/kg per d. Fig. 6(a) shows dopaminergic neurons in the SNpc. In the control group, many TH-positive cells were observed with normal shapes and neurite lengths, whereas in the MPTP group, few TH-positive cells were observed, and they had shrunken cell bodies and neurites. In contrast, the MPTP+ME group...
showed less dopaminergic neuronal cell loss than did the MPTP group. The immunoreactivity of TH-positive optical density in the ST showed a similar tendency (Fig. 6(b)). In the Bax immunostaining in the SNpc, Bax-immunopositive cells were increased in the MPTP group, whereas they were decreased in the MPTP + ME group (Fig. 6(i)–(k)).

Discussion

The present study showed that the ME protected dopaminergic cells from neurotoxicity in SH-SY5Y and primary mesencephalic cultures. Moreover, the present study confirmed the effect of ME in an MPTP-induced animal PD model, which included bradykinesia, nigral dopaminergic neuronal loss and striatal dopamine depletion. First, we used the MTT assay to investigate the protective effects of ME against 6-OHDA-induced neurotoxicity in SH-SY5Y cells. 6-OHDA, a neurotoxin which plays dominant neurotoxic roles in selectively damaging catecholaminergic neurons including dopaminergic neurons, has widely been used in experimental models of PD, and it can operate in extracellular or intracellular oxidation, yielding ROS that lead to toxic downstream molecules and resulting in neuronal damage(30). It has been demonstrated that 6-OHDA is involved with disturbing mitochondrial outer membrane permeability, leading to increased cytosolic cytochrome C and apoptotic proteins, including caspase-3(31). In the present study, ME showed a significant protective effect against 6-OHDA-induced toxicity and showed no toxicity to SH-SY5Y cells. In the measurement of ROS, ME significantly reduced intracellular ROS generation induced by 6-OHDA in SH-SY5Y cells in a dose-dependent manner. Because interaction between ROS and NO can stimulate the oxidative stress cascade and initiate a neurotoxic cascade(31,32), the inhibitory effect of ME on 6-OHDA-induced NO accumulation,

**Fig. 6.** Protective effect of ethanol extract of mulberry fruit (ME) on dopaminergic neurons in a mouse Parkinson's disease model. Saline or 500 mg/kg ME was administered orally once per day for 15 d, and 1-methyl-4-phenyl-1,2,3,6-tetrahydropyridine (MPTP; 30 mg/kg, intraperitoneally) was injected for the last 5 d. Dopaminergic neurons were visualised with tyrosine hydroxylase (TH) immunostaining. The numbers of TH-positive neurons in the substantia nigra pars compacta (SNpc) (a) were counted, and the optical density in the striatum (ST) (b) was measured. Representative photomicrographs of SNpc ((c)–(e)) and ST ((f)–(h)) were taken. Bax, an apoptotic protein, was visualised with Bax immunostaining in the SNpc ((i)–(k)). ((c), (f) and (i)) Control group; ((d), (g) and (j)) MPTP group and ((e), (h) and (k)) MPTP + ME group. Scale bar = 250 μm. Values are indicated as the mean values with their standard errors. *** Mean values were significantly different from the control group (P < 0.001). ††† Mean values were significantly different from the 6-MPTP-only-treated group (P < 0.001).
which was only about 10% but statistically significant, seems to contribute to the protective effect of ME.

Next, because ROS generated by 6-OHDA contribute to mitochondrial dysfunction, we measured Bax and Bcl-2 expression levels, $\Delta\psi_m$ and caspase-3 activation in SH-SY5Y cells to investigate the effect of ME on mitochondria-mediated apoptosis induced by 6-OHDA. Bac and Bcl-2 proteins have a role in apoptotic signal transduction by regulating the permeability of the mitochondrial membrane$^{[33]}$. $\Delta\psi_m$, an important factor in apoptosis, is disrupted by ROS, and it interacts with Bcl-2 family proteins$^{[34]}$. Caspase-3, which is produced by pro-apoptotic agent such as ROS and cascaded by 6-OHDA toxicity, appears to be essential in neuronal apoptosis through the apoptotic pathway common in many cell types$^{[35]}$. Present treatments for PD are focused on relieving symptoms; anti-apoptotic activity in a natural compound may improve the therapeutic options for treating PD and may be able to reduce the neurodegenerative progression$^{[36]}$. The present study showed that pre-treatment with ME regulated 6-OHDA-induced apoptotic pathway.

We also examined the effect of ME on GSH, an antioxidant protecting cells from toxins such as free radicals. 6-OHDA caused significant GSH depletion, whereas ME had no effect on GSH (data not shown). Thus, the neuroprotective effects of ME against 6-OHDA in SH-SY5Y cells may be mediated by ROS and NO inhibition, Bcl-2 and Bax regulation, $\Delta\psi_m$ stabilisation and inhibition of caspase-3 activation.

We also investigated the effects of ME on cultured primary mesencephalic dopaminergic neurons, which have been used to develop target agents for PD. MPP$,+,$ the active metabolite of MPTP, induces selective dopaminergic neuronal loss; it inhibits the respiratory complex I chain, leading to cell death$^{[36,37]}$. ME significantly protected dopaminergic neurons from 6-OHDA at a dose of 10 μg/mL. Against the other toxicity, MPP$,+,$ ME showed the effect at a wide range of concentrations of 10 and 100 μg/mL. From these results, we assumed that ME would be more effective in MPTP-induced mouse PD model than in other toxin-induced models.

Then, to evaluate the effect of ME in an in vivo PD model, we performed behavioural test and brain tissue stereology after MPTP and/or ME treatment. The MPTP mouse model is widely used for studying neuroprotective effects of candidate drugs because the characteristics of PD, such as oxidative stress, mitochondrial dysfunction and apoptosis, are reproduced pathologically and biochemically by MPTP$^{[36,37]}$. In the pole test, a commonly used behavioural test in the mouse model of PD$^{[38]}$, ME treatment, 500 mg/kg per d for 15 d, could maintain movement ability in MPTP-induced bradykinesia. In the histological analysis, whereas TH immunoreactivity was reduced in the SNpc and ST by MPTP, ME significantly reduced MPTP-induced dopaminergic neuronal damage in the SNpc and ST by inhibition of apoptotic protein, Bax. From these results, we confirmed that ME had neuroprotective effects in both in vitro and in vivo PD models.

In our previous studies, mulberry fruit showed a total phenolic content (TPC) of 1.78%, which is similar to that of the black bean, 1.82%, and more than that of red or white beans, 1.10% or 0.46%, respectively$^{[39]}$. Phenolic compounds are well known to have antioxidant effects, and they have been reported to inhibit 6-OHDA- and MPTP-induced apoptosis via the attenuation of oxidative stress$^{[40]}$. It has been reported that the value of oxygen radical absorbance capacity is correlated with phenolic contents in mulberry fruit, and that the total antioxidant capacity and TPC are comparable to those in other berries$^{[22,41]}$. Also, C3g, an aglycon of anthocyanin, has a neuroprotective effect against cerebral ischaemia via reduction of ROS generation$^{[21]}$. Polyphenols and other antioxidant compounds such as δ- and γ-tocopherols in ME$^{[22–24,27,28]}$ may have important roles in protecting against 6-OHDA- and MPTP-induced neurotoxicity via their anti-oxidant and anti-apoptotic effects, blocking of ROS and NO generation, regulating Bcl-2 family protein, stabilising mitochondrial membrane and inhibiting caspase-3 activity.

In summary, the mulberry fruit extract, by its antioxidant and anti-apoptotic effects, significantly protected neurons against neurotoxins in in vitro and in vivo PD models. These results suggest that mulberry fruit or compounds in it may provide neuroprotective candidates for use in treating or preventing PD.

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