The Protective Effects of Protein-Enriched Fraction from Housefly (Musca domestica) against Aged-Related Brain Aging

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Summary The Musca domestica larvae are well known for its multifunctions and great nutritional value. The present study aimed at investigating the beneficial effect of Musca domestica larvae extract (Mde) against memory impairment, structural damage and oxidative stress in aged rats. Twenty-month-old rats were gavaged with Mde for 2 mo. Morris Water Maze test indicated Mde prevented aging-induced spatial learning and memory dysfunction in the aged rats. Mde supply was also found to attenuate age-associated changes of brain histology that observed by light microscopy and transmission electron microscopy. Moreover, the increase of antioxidant capacity, glutathione peroxidase (GPx) activity, superoxide dismutase (SOD) activity, as well as the decreased methane dicarboxylic aldehyde (MDA) levels, were consistent with these results. Hence, we propose that oral administration of Mde could improve memory impairment via antioxidant action, and Mde has the potential to act as an excellent food supplement or medicine for the attenuation of brain aging.

Key Words Musca domestica larvae extract, antioxidants, aged rats, oxidative damage, memory impairment

The aged tendency of population is a major public health problem and one of the most complex issues in neuroscience today, which results in increasing public health concerns and health expenditures (1, 2). It is a widely accepted consensus that aging is typically accompanied by gradual and progressive cognitive decline and pathophysiological changes in the brain, which may cause age-related neurodegenerative diseases such as Alzheimer’s disease, Parkinson’s disease and so on (3, 4). The free radical (FR) theory was put forward to demonstrate the aged-associated cognitive decline and neuronal loss may be caused by the cumulative of oxidative damage in cells and tissues (5). Numerous studies also proved that the antioxidant supplements is an important step in the prevention of aging, because the antioxidants can scavenge free radicals or oxidation products that bring about oxidative damage (6, 7).

Insects and insert-derived products are extensive and unexploited sources of potentially useful compounds for modern medicine (3). Musca domestica larvae have been used as a therapeutic drug in China since Ming dynasty. Previous study has revealed that the extract of Musca domestica larvae has the ability to elevate antioxidant enzyme activities in vitro (8). Hence, it was expected that Musca domestica larvae extract may improve brain health against oxidative damage. Nevertheless, we rarely found its anti-aging aspect reports on age-related cognitive impairment in natural aging rats.

In the present study, the natural aging rat (20-mo-old) was used as a model to study the effect of Musca domestica larvae extract (Mde) on aged-related memory impairment. Structural alterations of brain were analyzed by optical microscope and transmission electron microscopy (TEM). The activities of relative antioxidant factors were measured by biochemistry and molecular biology methods.

MATERIALS AND METHODS

Preparation of protein-enriched extract from Musca domestica larvae. Musca domestica larvae were obtained from the Guangdong Provincial Center for Disease Control and Prevention (CDC), China. The protein-enriched extract of Musca domestica larvae was prepared as described before (9). Briefly, the third-instar larvae were collected, washed with distilled water, and homogenized with a homogenizer (5 s at 3,500 rpm). Next, the homogenate was centrifuged at 1,300 ×g/min for 10 min to remove insoluble material. The obtained supernatant was concentrated and lyophilized. Then the lyophilized supernatant, an extract of Musca domes-
tica larvae (Mde), was stored at −80°C (10). The nutritional value of Mde has been determined in our previous studies (9, 11).

**Animals and experimental design.** All animal experiments were carried out according to the guidelines of the 3Rs (Replacement, Reduction and Refinement) after approval by the Ethical Committee for Research on Laboratory Animals of Guangdong Pharmaceutical University [China approval number SCXK (Yue) 2013-0002]. Adult female Sprague-Dawley (SD) rats (3 mo, 220 to 250 g) and natural aging female SD rats (20 mo, 350 to 450 g) were procured from Medical Laboratory Animal Center (Guangzhou, Guangdong, China). All rats were kept in specified pathogen free (SPF) conditions and maintained on a light-dark cycle (12 h/12 h) at 25±2°C with 55±10% relative humidity.

The rats were fed ad libitum with normal diet for 1 wk. Rats were fed with food provided by the Animal Center of Guangdong Pharmaceutical University (protein 22%, fat 4.2%, calcium 1.2%, phosphorus 0.9%, lysine 1.36%, protein+cystine 0.82%). After acclimatization, eight 3-mo-old SD rats were served as young control group. Forty 20-mo-old rats were randomly assigned to five groups: aged control group (normal saline, n=8), vitamin E (VitE) group (n=8), the low-dose Mde group (50 mg/kg, n=8), the middle-dose Mde group (100 mg/kg, n=8) and the high-dose Mde group (200 mg/kg, n=8). All rats were given intragastric administration once a day at the same time, at the dose of 10 mL/kg for 2 mo.

All rats were tested in Morris Water Maze after 2 mo. After the behavior test, all animals were deeply anesthetized with 2% sodium pentobarbital. For ultrastructural study, some rats’ thoraces were opened, isotonic saline was quickly perfused into the aorta via left ventricle, and then ice cold 2.5% glutaraldehyde that had been diluted in 0.1 M phosphate buffer were slowly perfused in the same way. Following perfusion, their brains were taken out immediately. In addition, the others’ brains were rapidly removed, washed with ice-cold normal saline and stored at −80°C till further analysis.

**Behavioral study—Morris Water Maze (MWM).** Behavior study was performed by Morris Water Maze test to observe the spatial learning and memory of rats (12). The test apparatus consisted of hidden escape platform (a 10 cm diameter round iron) and round pool (1.20 m diameter, 60 cm height), filled with water in the depth of 40 cm with temperature of 25±1°C. Four equidistant points were designated as start positions and divided the pool into four quadrants. The hidden escape platform was placed under the water surface 2 cm in the center of southwest quadrant. Ink was poured into the water to make water turbid to prevent extra-maze cues (e.g. laboratory technician, table, window, cabinets, furniture and light). The test was carried out in a dark and sound proof room.

At the end of 60th days, rats were put into the pool 1 min before starting the experiment to allow habituation to the new environs. Behavior analysis was performed twice daily, lasting for 3 d. The experimenter did not know the group of rats in the course of the experiment, so as to avoid the influence of subjective consciousness guidance on the experimental results. At the beginning of each test, rats were placed in the pool, facing each wall of the four quadrant edges in a pseudo-random manner. The rats were allowed to find the hidden escape platform in 60 s, or they were gently guided to the platform by the experimenter. Then the rats were given a 10 s rest on the platform in the test interval time. The behavioral data were collected through a video camera fixed on ceiling and analyzed by the MNT-100 Video Tracking System (Chengdu Tai meng Tech. Co, Ltd, Chengdu, China).

**Hematoxylin–eosin staining (HE).** For histological analysis, brains were fixed with 4% paraformaldehyde for at least 24 h, dehydrated in a series graded alcohols and embedded in paraffin. After which serial, 4 µm slices were cut and transferred onto pre-treated glass slides. Sections were deparaffinized and stained with hematoxylin and eosin for examining the morphological changes in the brain. Sections were selected from all animals at the same level of the coronal section, one section was collected every 5 consecutive sections until a total of 5 sections had been acquired for each animal. Five visual fields (×400) were randomly selected to count the numbers of preserved nerve cells per 1 mm² in the cortical regions in a single-blinded manner by the Image J software (National Institutes of Health, Bethesda, MD, USA) (13).

**Transmission electron microscopy.** For brain ultrastructure observing, the brain were diced into blocks less than 1 mm³ and fixed in 2.5% glutaraldehyde. After further postfixation in fresh fixative (1% OsO₄) for 2 h with the temperature of 4°C, the blocks were dehydrated in ascending series of alcohols and then embedded in epoxy resin for preparation of slices. The 0.5 mm thick slices were stained with 1% Toluidene blue and observed in an optical microscopy for further repairing the embedded block. Ultraline sections were obtained by the use of an ultramicrotome with 1 mm/s speed and 100 nm thickness. Finally, the ultrathin section were placed on a mesh with supporting film before staining Urynl Acetate-Lead Citrate stain in a petri dish, to absorb the carbon dioxide at room temperature for 3 min, the grid was removed and the slices were observed in a JEM1400 (Jeol, Tokio, Japan) transmission electron microscope (14, 15). Micrographs were taken with a digital camera Bioscan 792 (Gatan Inc, Pleasanton, USA).

**Measurement of biochemical indexes.** The brains were homogenized by an organization dispersion machine (IKA, Germany). Next the homogenates were centrifuged at 3,000 rpm/min for 15 min. Supernatant was collected for the measurement of oxidative stress indicators such as superoxide dismutase (SOD), glutathione peroxidase (GPx) and methane dicarboxylic aldehyde (MDA). The level of SOD, GPx and MDA of rats in the brain homogenate was detected by the using the assay kits in accordance with specification requirements (Nanjing Jiancheng Bioengineering Institute, Nanjing,
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The naturally-induced aging rats with and without the treatment of Mde was evaluated by Morris Water Maze. (A) The swimming distance of all groups to locate the hidden platform over 3 d. (B) The escape latency of all groups to locate the hidden platform over 3 d. (C) The swimming speed of all groups to locate the hidden platform over 3 d. Data are represented as mean±SD. Compared with the aged control group, *p<0.05, ***p<0.001.

**RESULTS**

**Behavioral alterations**

The behavioral analysis is a sensitive method for revealing the impairment of spatial learning and memory (20). The swimming distance, latency and speed to locate the hidden platform of all rats are shown in Fig. 1. The test showed that the aged rats took longer to find the hidden platform in second and third days compared with young control group (p<0.05), suggesting that the normal aging rats had significantly cognitive impairment. We found that the prolonged swimming distance and escape latency in the aged control group were shortened by administration of Mde or VitE especially on second day (Fig. 1A, B). Furthermore, the swimming speed of rats in the aged control group was slower than that of the other group (Fig. 1C), but no significant difference was observed either between the aged group and the Mde-treated group or between the aged and young control group (p>0.05). Considered together, these results indicated that Mde prevented aging-induced spatial learning and memory dysfunction in rats, and the effect of Mde was not less than that of the more well-known VitE.

**Hematoxylin–eosin staining**

In the young control group, the neurons had normal and clear morphological structure. Cortical cells were arranged in high density (Fig. 2A). The cerebral cortex of the aging rats showed pronounced pathological changes compared to the young rats. A higher number of cells were densely stained, which exhibited pyknotic nuclei and shrinkage of cytoplasm. Additionally, irregularly shaped cells also were seen (Fig. 2B). However, the shrinkage of the neurons was rarely found in the aging rats that treated with high dose of Mde (Fig. 2F). Compared with adult rats, the ratio of neurons in cerebral...
Fig. 2. Representative HE images of cerebral cortex. (A) young control group, (B) aged control group, (C) aged rats treated with VitE, (D) aged rats treated with low-dose Mde, (E) aged rats treated with middle-dose Mde, (F) aged rats treated with high-dose Mde.

Table 1. The quantitative statistics of neuronal parameters in cerebral cortex.

| Group | Neuron-density/mm² | Cell-density/mm² | Ratio of neurons/% | Ratio of damaged neurons/% | Max-diameter of neuronal nuclear/µm |
|-------|-------------------|------------------|--------------------|---------------------------|----------------------------------|
| A     | 107.67±42.03      | 4,226.11±107.67  | 2.55               | 0                         | 13.58                            |
| B     | 74.02±25.77       | 3,425.30±74.02   | 2.16               | 0.982                     | 10.18                            |
| C     | 60.57±20.19       | 2,839.84±60.57   | 2.13               | 0.711                     | 11.04                            |
| D     | 67.29±11.66       | 3,297.44±67.29   | 2.04               | 0.408                     | 10.93                            |
| E     | 80.75±38.07       | 3,930.01±80.75   | 2.05               | 0.514                     | 10.86                            |
| F     | 67.29±34.75       | 3,082.10±67.29   | 2.18               | 0                         | 12.75                            |

(A) young control group, (B) aged control group, (C) aged rats treated with VitE, (D) aged rats treated with low-dose Mde, (E) aged rats treated with middle-dose Mde, (F) aged rats treated with high-dose Mde. The main criteria for judging neurons included large cell body, large and round nucleus (diameter >6.5 µm), shallow staining and obvious nucleolus. The main criteria of damaged neurons included dense staining, pyknotic nuclei and shrinkage of cytoplasm.

Fig. 3. TEM images of cerebrum neurons after the experiment. (A) young control group, (B) aged control group, (C) aged rats treated with VitE, (D) aged rats treated with low-dose Mde, (E) aged rats treated with middle-dose Mde, (F) aged rats treated with high-dose Mde. Yellow arrows point to the lysosome. Red arrows point to the intracellular mitochondria. Purple arrows point to the extracellular mitochondria. Blue arrows point to the rough endoplasmic reticulum. Green arrows point to the synapses.
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The cortex of aging rats has not changed much. Increased proportion of damaged neurons and smaller diameter of neurons was found in aging rats (Table 1). Pathological damage impaired by aging could be prevented from the Mde or VitE and the effect of Mde appeared in a dose-dependent manner (Fig. 2D–E).

**Transmission electron microscopy**

In the young control group, degenerating neurons were rarely found. The neurons showed intact nuclear with evenly distributed chromatin. The cytoplasm contained intact organelles, such as well-developed mitochondria and rough endocyttoplasmic reticulum with rosettes of polyribosomes (Fig. 3A). The synapses of neurons were intact and smooth (Fig. 4A). In the aged control group, neurons had some iconic aging characteristics, consisting of cytoplasmic vacuolation and degenerative changes in the cytoplasmic organelle. Neurons also showed a decrease in their synaptic coverage compared to young controls. Furthermore, endoplasmic reticulum was dilated, fragmented or vacuolated. Ribosomes were obviously detached from the external surface of the rough endoplasmic reticulum. Mitochondrial were swollen without visible cristae. Synaptic gap junction became less and fuzzy. Lots of lysosomes appeared in the cytoplasm of neurons (Figs. 3B, 4B). With the treatment of Mde, the pathological changes had been evidently improved (Fig. 3C–E; Fig. 4C–E). For example, mitochondrial structure tends to be normal. Lysosomes had fewer numbers in the cytoplasm. The number of neural connections increases. The coloration of synaptosomes was darker. These findings indicated Mde can improve the pathological state of neurons and delay the process of cerebrum senility.

**Measurement of biochemical indexes**

The oxidative stress proteins (SOD, GPx, MDA) after perfusion with Mde as compared with controls was detected. The SOD and GPx activities in aged control group were found to be lower than young control group (Fig. 5A, C). Compared with the aged control group, the SOD and GPx activities of all Mde-treated rats or VitE-treated rats were found to be significantly increased ($p < 0.05$). Compared with young control group, the MDA content was also easily found to be significantly increased in aged control group (Fig. 5B). Interestingly, Mde or VitE can considerably down-regulate MDA levels in aging rats ($p < 0.05$). In the present study, SOD and GPx activities increased dose-dependently, and the level of MDA decreased in the Mde-treated groups. This phenomenon demonstrates that antioxidant activity is improved and markers of oxidative stress are diminished in Mde-treated aging rats.

**PCR, QPCR and IHC**

The mRNA content of SOD2 in natural aging rats was significantly decreased compared to young rats (Fig. 6A, B). The expression of SOD2 in aging brain was significantly increased after treatment with Mde ($p < 0.001$). The aging rats with the treatment of Mde reversed the level of SOD2 near to the young rats.

Brown positive particles were predominantly in the

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**Fig. 4. TEM images of neurons after the experiment.**

(A) young control group, (B) aged control group, (C) aged rats treated with VitE, (D) aged rats treated with low-dose Mde, (E) aged rats treated with middle-dose Mde, (F) aged rats treated with high-dose Mde. Purple arrows point to the extracellular mitochondria. Green arrows point to the synapses.

**Fig. 5.** The SOD, MDA and GPx activity was observed by several biochemical methods. (A) The activity of SOD in the brain. (B) The content of MDA in the brain. (C) The activity of GPx in the brain. Data are represented as mean ± SD. Other groups were compared with the aged control group, ∗$p < 0.05$, ∗∗$p < 0.01$, ∗∗∗$p < 0.001$. 

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**Fig. 6.** The expression of SOD2 was examined by QPCR in brain. The expression of SOD2 was significantly increased after treatment with Mde ($p < 0.001$). The aging rats with the treatment of Mde reversed the level of SOD2 near to the young rats.
granular cell layer (GCL) and polymorphic layer (PL). The positive rate of SOD2 was reduced by about 78.27% in natural aging rats than in young rats. Compared with aged control group, the positive rate of SOD2 was increased by about 65.76% after perfusion with Mde (Fig. 6C). These phenomena indicated that Mde induce the increase of antioxidant gene and protein.

**DISCUSSION**

Natural aging rats are widely used for studying age-related pathogenesis in domestic and foreign laboratory (21). Memory decline is a common phenomenon in the aging process (22, 23). The prolongation of escape latency in aged rats in the Morris Water Maze test is thought to reflect age-related learning and memory impairment (24–26). After administration of Mde, the learning and memory of aging rats were significantly improved as evident from the reduced escape latency and swimming distance compared to the aged control group. This is the first report about the improvement of Mde on memory impairment in natural aging rat models.

The cerebral cortex that contains countless nerves and cells controls the minds and actions of the human body. It has been found that aging can reduce the subsets of cerebral neurons and inhibit the proliferation of neurons through immune T cells (27, 28). HE staining revealed that the neuronal cells in cerebral cortex were significantly lost and atrophied, and the proportion of damaged neurons increased as compared to adult rats. The changes of neuronal cell density and architecture of neurons were pathological modifications that might happen in the brain in aging processes (29). In addition, the diameter of neurons tends to decrease, and the neuronal cell is shrinking rather than disrupt, which is probably a more decisive age-related alteration (30). It is speculated that Mde can reduce the injury of neurons and is beneficial to the normal growth of neurons. In our study, oral administration of Mde showed a neuroprotective response with significantly attenuated the shrinkage of the neurons and improved organization of cellular layers.

To further investigate the changes of brain ultrastructure, we used the transmission electron microscopy to observe the temporal lobe of neurons. The aging rats exhibited pathological changes including irregularly nuclei in shape and cytoplasmic vacuolization. Furthermore, organelle showed degenerative changes, including lysosomal expansion, mitochondrial edema, endoplasmic reticulum expansion, ribosome shedding and synaptic space stenosis. Decreased cellular function is associated with changes in cell structure. Mitochondrial damage could lead to a decline in hypoxia tolerance, oxidative phosphorylation reaction and protein synthesis. The endoplasmic reticulum and ribosome damage will result in low protein synthesis. The decrease in protein synthesis will affect the cell structure and metabolic processes as well as structural protein and enzyme synthesis. Lysosome can remove aging organelles and useless biological macromolecules. The presence of lysosome is one sign of cellular senescence. Signal transduction between neurons relies on nerve synapses to complete. The abnormalities of information exchange between neurons is closely associated with age-related decline (31). Interestingly, Mde supply was found to attenuate age-associated changes of brain histology and ultrastructure. The structural changes proved Mde has beneficial effect against structural damage in aged brain.

The accumulation of oxygen-centered free radicals (FRs) and other reactive oxygen species (ROS) during aging may contribute to the oxidative damage, which can lead to several events such as loss in protein synthesis, depletion of cellular redox balance and ultimately to
cell aging and death (32). SOD is the primary substance of free radical scavenging in the organism (33, 34). The level of SOD in vivo is an intuitive index about aging and mortality. Glutathione peroxidase (GPx) is an important peroxidative enzyme, which can protect the structure and function of cell membrane from peroxides. Lipid peroxidation end products such as MDA can induce the crosslinking polymerization of macromolecules. Aging rats is characterized by the decreased activity of antioxidant enzymes and increased lipid peroxidation (31). With the treatment of Mde, the antioxidant activity was improved and markers of oxidative stress were diminished in aging rats. These finding indicated that the protection of Mde against age-related neurons damage may be reversed by antioxidants. Moreover, SOD2 display an ability to preserve proliferation of neural stem cells in the aging rats (35). The expression of SOD2 mRNA and protein in brain were examined using PCR, QPCR and immunofluorescence analysis. The results indicated that increased SOD2 in the cerebral cortex may protect the neurons from superoxide-induced damage in Mde-treated aging rats. These results indicated that the protection of Musca domestica larvae extract against age-related memory impairment may be related to attenuation of oxidizing free radical-generated oxidative stress.

It is reported that Musca domestica larvae extract (Mde), the protein-enriched extract of Musca domestica larvae, have significant antioxidant activity (8). In the present study, we found the supplementation of Mde may protect brain from age-related neuronal loss and memory impairment by reducing oxidative stress. The excessive free radical oxidation may damage the normal function of neurons, and ultimately evolve into neurodegenerative diseases (6, 7). Brain protection may provide new ideas and application prospects for neurodegenerative diseases (36). In view of the different chemical compositions of VitE and Mde, an appropriate positive control must be necessary in the future studies, such as Tenebrio molitor or Silkworm pupae extract, which contains nutritional value similar to Mde (37, 38) and could be better used as a control for delineating specific biological actions of Mde to aged brain. Then, further study need to be carried out to explore the role of Mde in aging-related diseases.

In conclusion, we provide the evidence that the Mde could improve memory impairment and help neurons to maintain normal physiological structure in aging brain. The protective effects of Mde may be realized by reducing oxidative stress and increasing antioxidant enzymes. Musca domestica larvae extract may act as a dietary supplement or developing a medicine to delay brain aging.

Authorship
Research conception and design: Yanan Tang and Xuemei Lu; experiments: Yanan Tang, Panpan Feng, Shuqing Gui, and Xuemei Lu; statistical analysis of the data: Xiaobao Jin and Jiayong Zhu; interpretation of the data: Yanan Tang, Panpan Feng and Xuemei Lu; writing of the manuscript: Yanan Tang and Xuemei Lu. All authors discussed the results for completion of the manuscript.

Yanan Tang, Panpan Feng and Shuqing Gui had equal contribution to this article.

Disclosure of state of COI
No conflicts of interest to be declared.

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Supporting information
Supplemental online material is available on J-STAGE.

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