Therapeutic role of bone marrow mesenchymal stem cells in diabetic neuronal alternations of rat hippocampus

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Background: As the hippocampus is the main brain region for many forms of learning and memory functions and is acutely sensitive to blood glucose changes, diabetes mellitus, which is a serious metabolic disease, is often accompanied by learning and memory deficits. Through scientific literatures, mesenchymal stem cells (MSCs) promote functional recovery in rats with traumatic brain injury, so the present work was conducted to study MSCs as a possible treatment for the diabetic neuronal degeneration and functional impairment of rat hippocampus.

Materials and methods: It was carried out using male albino rats: non-diabetic control groups (4, 8, 12 weeks) (n = 15), diabetic groups by i.v. injection of streptozotocin for (4, 8, 12 weeks) (n = 15) and MSCs treatment to diabetic groups for (8, 12 weeks) (n = 10). Hippocampal learning and memory functions were assessed by the Morris Water Maze test and its results were statistically analysed. The rat hippocampal regions (CA1 and CA3) were subjected to histological, ultrastructural examination and morphometrical analyse of pyramidal neurons.

Results: Neurons of the diabetic groups showed disturbed function and architecture; shrunken hyperchromatic nuclei and vacuolated eosinophilic cytoplasm (apoptotic changes) also MSCs treatment improved hippocampal learning and memory functions plus its architectural changes; increasing populations and normal regular distribution.

Conclusions: It can be concluded that diabetic hippocampal neuronal alternations and functional impairment can be ameliorated by MSCs treatment. (Folia Morphol 2020; 79, 2: 211–218)

Key words: diabetes, mesenchymal stem cells, hippocampus, neuronal alternations, learning and memory

INTRODUCTION

Uncontrolled diabetes mellitus (DM) leads to severe complications of central nervous system (CNS) usually associated with neuronal degeneration [31]. As the hippocampus is particularly sensitive to changes in blood glucose level specially CA1 and CA3 regions [21], DM produces hippocampal dysfunction that is involved in learning and memory processing enhancing the risk of Alzheimer’s disease [3]. Through scientific literature, DM can affect the hippocampus through reducing its synaptic plasticity [22], changes glutamate neurotransmission [10] and augments enzymatic activity linked to
oxidative stress in the hippocampus [29]. These changes are usually associated with signs of neuronal loss and apoptosis which are due to stimulation of caspase-3 activity, nuclear DNA cleavage and induction of proapoptotic genes in hippocampus especially with uncontrolled DM [19]. As streptozotocin (STZ) can cause pancreatic β-cell destruction, it is used experimentally to induce type 1 DM [17]. Mesenchymal stem cells (MSCs) have been used experimentally in the treatment of the injured brain [1, 6, 11] as well as other damaged organs as in kidney [9], heart, etc. In the reviewed literatures there was obvious contradiction concerning the differentiation of MSCs into the functioning nerve cells. Several studies showed that MSCs transplanted into the intact, injured or diseased CNS environments do not differentiate or even a small portion of them produce neural phenotypes [8]. Even though, Chopp and Li [6] demonstrated that, transplanted MSCs promote functional recovery in rats with traumatic brain injury through activation of endogenous angiogenesis, neurogenesis, and synaptogenesis. Moreover, other studies supported that MSCs could differentiate into mature neuron-like cells and exhibit neuronal properties [1, 11]. Also, Calió et al. [4] discussed MSCs role on the injured brain by decreasing neuronal apoptosis and oxidative stress. The aim of the present work was to evaluate using MSCs as a possible treatment for diabetic neuronal degeneration and learning and memory alternations.

MATERIALS AND METHODS

MSCs preparation

Mesenchymal stem cells were obtained from Medical Biochemistry Department, Faculty of Medicine, Cairo University. Bone marrow derived MSCs was isolated and cultivated for 4 weeks according to protocol of Jiang et al. [13]. Next, cells were labelled with 5-bromo-2′-deoxyuridine (BrdU). Fluorescence phase-contrast microscope (Axiocam MR R3, Carl Zeiss, Germany) was used to observe the rat MSCs every 2 or 3 days.

Animals

The experiment strictly adhered to all ethical guidelines regarding animal research and was approved by the Institutional Animal Care and Use Committee of Cairo University (CU-IACUC). Adult normoglycaemic male Sprague-Dawley albino rats (150–200 g) were used in the current study. Rats were kept in a temperature- and humidity-controlled room, with free access to food and water and were placed 5/cadge.

Experimental design

The rats were divided into three groups.

Control group (n = 15) which was subdivided into three subgroups (n = 5/subgroup) [20]: (Control, 4 weeks: sacrificed after 4 weeks, Control, 8 weeks: sacrificed after 8 weeks, and Control, 12 weeks: sacrificed after 12 weeks).

DM groups (n = 15) received single i.v. injection of STZ (STZ, Sigma–Aldrich, St. Louis, MO, USA) dissolved in sodium citrate 0.1 mL buffer, PH 4.5 at a dose of 60 mg/kg [2]. This group was subdivided into three subgroups (n = 5/subgroup) [20]: (4-week DM: sacrificed after 4 weeks, 8-week DM: sacrificed after 8 weeks, and 12-week DM: sacrificed after 12 weeks). Fasting blood glucose levels were measured in DM groups (using spectrophotometer-Beckman; USA) 72 h after STZ injection to confirm induction of DM (> 200 mg/dL). This day was considered the first day of the experiment.

DM + MSCs group (n = 10) received single i.v. dose of STZ and 4 weeks later a single i.v. dose of fluorescent MSCs 1 mL of about 2,000,000 stem cells in the rat tail vein [15]. This group was subdivided into two subgroups (n = 5/subgroup) [20]: (8-week DM + MSCs: sacrificed after 8 weeks from the start of experiment; 4 weeks after MSCs administration and 12-weeks DM + MSCs: sacrificed after 12 weeks from the start of experiment; 8 weeks after MSCs administration). Fasting blood glucose levels were measured in DM and MSCs groups before sacrification.

Morris Water Maze (MWM) test

Learning and memory impairment induced by DM and improvement after MSCs treatment were evaluated through performing MWM (4, 8 and 12 weeks from the start of the experiment). It was performed according to previous studies by Vorhees and Williams [32]. Four-day training (twice/day) learning and memory was assessed through measuring escape latency to rich hidden platform (recorded with video camera). The maze was built of white circle tank (diameter: 127 cm and height: 51 cm) filled with water depth 45 cm (made opaque with liquid milk) and a white platform (12 cm²) submerged 2 cm below the water surface in the northwest quadrant of the maze. Navigational landmarks in the form of external cues were placed to the room wall. Signs of retrieving such memory (by removing platform and animals were allowed to swim freely for 60 s) were assessed through a provisional trial by counting both the number of platform-site crossovers and percent-
age of time spent in the target quadrant compared with the other quadrants. Statistical analysis was done for the means of escape latency times/secs of the trails and for the provisional trial.

**Sacrification and histological staining**

By the end of the experimental periods, animals were anaesthetised by i.v. injection of ketamine-dylazine and cardiac perfusion fixation technique by formalin 10% was done. For brain extraction, animals were decapitated immediately rostral to the first cervical vertebra and brains were removed. The right hemispheres allocated for light microscopic examination stained with haematoxylin and eosin (H&E). Studying hippocampal both CA1 and CA3 regions, as those regions are the hub of memory and learning functions [18]. The left hippocampi were dissected and allocated for ultrastructural examination using transmission electron microscope JEOL TEM (JEM-1400, JEOL, Japan) at 80 kV. To detect labelled MSCs in the hippocampus, 2 sections of both MSCs subgroups were immunostained by anti-Brdu and goat anti-mouse Ig and examined by fluorescent microscopy (Fig. 1).

**Morphometric analysis**

It was conducted by using hippocampal histological sections stained by H&E, implemented for the numbers of the pyramidal neurons in CA1 and CA3 hippocampal regions in all subgroups (5 sections par each) using total magnification of ×400. These statistical assessments were performed using analysis of variance (ANOVA) and statistical package for social science (SPSS). P value > 0.05 was considered not significant (NS) and p value < 0.05 was considered significant (S).

**RESULTS**

**Statistical analysis**

Regarding to the escape latency parameter in MWM test, all control groups showed identical numerical data so they were considered as one control group. All DM subgroups tended to require significant more escape latency periods to find platform than control group and there was more significant difference for 12-week DM subgroup than 8-week DM subgroup (p < 0.05) (Fig. 2A). On the other hand, MSCs treatment improved learning and memory deficit, as both MSCs subgroups exhibited a significant decrease in the escape latency period (p < 0.05) as compared to DM subgroups (Fig. 2A) but still significantly higher numbers when compared to control groups (p < 0.05) (Fig. 2A). Evaluation of provisional trial test showed the mean values of the DM subgroups were significantly lower compared to control groups (p < 0.05). However, the outcomes of DM + MSCs subgroups were significantly improved when compared to DM subgroups, but still lower than that of control groups (p < 0.05) (Fig. 2B).

Concerning fasting blood glucose levels in all groups, the mean values were significantly increased in DM groups according to the extent of diabetic durations (p < 0.05) but with MSCs treatment the values were significantly decreased according to the extent of treatment indicating improvement (p < 0.05) (Table 1).
Regarding to the morphometrical statistical analyses of the pyramidal neurons, mean values were used to assess diabetic neuronal degeneration and regeneration by MSCs of CA1 and CA3 regions. The numbers of both regions of DM subgroups were decreased when compared to control group. Pyramidal cell reduction of CA3 was significant in 8-week and 12-week DM subgroups while the reduction of number of cells of CA1 was only significant in 12-week DM subgroup (p < 0.05). Moreover, among DM subgroups, their mean numbers were decreased according to the extent of DM duration; this reduction was significant in rats of 12-week DM subgroup (p < 0.05). There was prominent neuronal number improvement in DM + MSCs subgroups. However, the values were still significantly lower in 8-week DM + MSCs subgroup when compared to control group (p < 0.05). On the other hand, marked improvement in their numbers was demonstrated in rats of 12-week DM + MSCs subgroup which was non-significant in CA1 region and only significant in CA3 region when compared to the control group (p > 0.05) (Fig. 3).

Haematoxylin and eosin stained sections

All control groups showed identical histological architecture, so they were considered as one control group.

Examination of CA1 region. The control group revealed arranged pyramidal neurons of uniform size and shape with narrow neuropil in-between, each en- closing single, rounded, large and vesicular nucleus with prominent nucleolus. Many glial cells within the molecular layer were seen (Fig. 4A). On examination of 4-week DM subgroup, most of the pyramidal neurons were comparable to the control, while other few neurons were shrunked with karyorrhectic and karyolytic nuclei. Vacuolations were also seen (Fig. 4B). On examination of the 8-week DM subgroup showed more neuronal damage as many shrunken nuclei (arrows) and clumping neuronal processes (arrowheads); C. Eight-week DM: clogged margined chromatin (arrows) and hyperchromatic neurons (curved arrows) in molecular layer (M); D. Twelve-week DM: severe degeneration as, many shrunken nuclei (arrows) and clumping neuronal processes (arrowheads); E. Eight-week DM + MSCs: neuronal improvement (thick arrows), few shrunken nuclei (curved arrows) and clumping processes (asterisk); F. Twelve-week DM + MSCs: progressive neuronal improvement (thick arrows), few shrunken neurons (thin arrows); bar = 50 μm; DM — diabetes mellitus; MSCs — mesenchymal stem cells.

Table 1. The mean values of fasting blood glucose level (mg/dL) in all groups

<table>
<thead>
<tr>
<th>Control groups</th>
<th>DM, 4 w</th>
<th>DM, 8 w</th>
<th>DM, 12 w</th>
<th>DM + MSCs, 8 w</th>
<th>DM + MSCs, 12 w</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control</td>
<td>82.7 ± 5.65</td>
<td>224 ± 6.91</td>
<td>248.5 ± 35.08</td>
<td>273 ± 13.93</td>
<td>134.5 ± 22.59</td>
</tr>
</tbody>
</table>

DM — diabetes mellitus; MSCs — mesenchymal stem cells; w — weeks

Figure 4. CA1 region; A. Control group: arranged pyramidal (P) with vesicular nucleus (thick arrows) and glial cells (thin arrows) in molecular layer (M); B. Four-week DM: karyorrhectic (thick arrows), karyolytic (zigzag arrow) and shrinkage nuclei (thin arrows), vacuolations (arrowheads); C. Eight-week DM: clogged margined chromatin (arrows) and hyperchromatic neurons (curved arrows) in molecular layer (M); D. Twelve-week DM: severe degeneration as, many shrunken nuclei (arrows) and clumping neuronal processes (arrowheads); E. Eight-week DM + MSCs: neuronal improvement (thick arrows), few shrunken nuclei (curved arrows) and clumping processes (asterisk); F. Twelve-week DM + MSCs: progressive neuronal improvement (thick arrows), few shrunken neurons (thin arrows); bar = 50 μm; DM — diabetes mellitus; MSCs — mesenchymal stem cells.

Figure 3. The mean numbers of the pyramidal neurons in regions (CA1 and CA3) in all subgroups; w — weeks.
mic Nissl granules surrounded by narrow neuropil. The molecular layer showed glial cells (Fig. 5A). However in 4-week DM subgroup, the pyramidal neurons represented normal nuclear appearance, while few nuclei with clogged marginated chromatin, relatively few shrunken neurons with vacuolated cytoplasm or with peripheral karyorrhectic nuclei and narrow neuropil with few vacuolations (Fig. 5B). Moreover, on examination of 8-week DM subgroup there were pleopathological changes, such as pyramidal neurons haphazardly arranged with many pyknotic nuclei, neurons with peripheral homogenous nucleus and severe vacuolation with clumping processes (Fig. 5C). Also, the 12-week DM subgroup showed more degeneration reflected by neuronal disorganisation with eosinophilic vacuolated cytoplasm and severe clumping processes (Fig. 5D). While on examination of 8-week DM + MSCs subgroup, there was minimal regeneration: pyramidal neurons showed normal appearance and almost normal neuropil comparable to control group. However, others were represented with karyolytic or pyknotic nuclei and vacuolated cytoplasm. Extraneuron amyloid deposition was noticed (Fig. 5E). Additionally, 12-week DM + MSCs subgroup had regeneration signs as most of the pyramidal neurons were comparable to control with vesicular nuclei. Only few shrunken neurons represented with hyperchromatic nucleus and areas devoid of neurons were seen (Fig. 5F).

**Ultrastructural study of the pyramidal neurons**

The control groups revealed the normal neuronal ultrastructure: intact cell membrane, intact cytoplasmic mitochondria with normal cristae. Euchromatic nucleus with smooth nuclear envelope consisting of bilamellar layers and well-formed nuclear pores were seen (Fig. 6A). On the other hand, the pyramidal neurons of 4-week DM subgroup showed dense neuron form with irregular outer line, ballooning mitochondria with disrupted cristae and cytoplasmic deposition of electron-dense bodies. The nucleus showed rounded and distinctly chromatin clumps (Fig. 6B). Pyramidal neurons of 8-week DM subgroup had more degeneration, such as nuclear condensation, ballooned mitochondria and more cytoplasmic lipofuscin particles (Fig. 6C). Additionally, pyramidal neurons of 12-week DM subgroup had severe degeneration signs...
and apoptotic changes, such as nuclear membrane indentation, cytoplasmic condensation, few neurons with highly condensed dense chromatin and vacuolations within the neuropil (Fig. 6D). While on examination of pyramidal neurons of 8-week DM + MSCs subgroup, there were some evidence of regeneration, such as normal neuronal form with euchromatic, central, and rounded nuclei. But few cells had abnormal dense form with shrunken and electron dense nuclei (Fig. 6E). Moreover, in pyramidal neurons of 12-week DM + MSCs subgroup, there was more evidence of regeneration, such as normal neuronal ultrastructure with intact mitochondria (Fig. 6F).

**DISCUSSION**

Diabetes mellitus is a common serious metabolic disease and usually associated with cerebral alterations [31]. Several studies indicated that DM negatively affects hippocampal cellular morphology, proliferation and survival, indicating function impairment of this brain region [35]. MSCs treatment had been shown as a promising modality in the treatment of the injured brain [11], beside that there were contradictions concerning the differentiation of MSCs into the functioning nerve cells. In the present study, the MWM results of DM subgroups indicating learning and memory impairment which become worst with prolonged durations. Similar results were obtained by Malone et al. [23] who further explained that DM decreased hippocampal neurotransmitters’ release: presynaptic synaptophysin and postsynaptic density protein, which is the cause of memory and learning deficiency. However in MSCs treated subgroups, there was prominent functional improvement which become worst with prolonged durations. These results were obtained by Malone et al. [23] who further explained that DM decreased hippocampal neurotransmitters’ release: presynaptic synaptophysin and postsynaptic density protein, which is the cause of memory and learning deficiency. However in MSCs treated subgroups, there was prominent functional improvement which become worst with prolonged durations. These results were obtained by Malone et al. [23] who further explained that DM decreased hippocampal neurotransmitters’ release: presynaptic synaptophysin and postsynaptic density protein, which is the cause of memory and learning deficiency.

Alongside with these functional impairment and improvements, the histological and ultrastructural examination showed hippocampal degenerative changes accompanying DM were variable regarding the affected region and also related to the duration of DM exposure. Minor apoptotic alternations were detected in 4-week DM subgroup in both CA1 and CA3 regions, in agreement with findings Zhao et al. [36] who study CA1 and dentate gyrus. However, more changes were mainly seen in CA3 region of 8-week DM subgroup concurring to findings of Jafari et al. [12] and Kamal et al. [14]. Furthermore, when DM was prolonged for 12 weeks, progressive changes were demonstrated mainly in CA3 region. These results were in harmony with findings of long-term study of Yang et al. [33].

These apoptotic degenerative changes were confirmed by more accurate statistical morphometric data as all DM subgroups showed reduction of number of pyramidal neurons which was more significant at 12 weeks than 4 weeks and 8 weeks indicating the progressive degenerative changes.

The findings of earlier diabetic neuronal apoptotic changes detected in the current work were explained by Zhao et al. [36] and Orlovsky et al. [27] as excessive production of nitric oxide in the hippocampal areas which has neurotoxic effect inducing neuronal damage and apoptosis. Also, the findings of progressive diabetic neuronal apoptotic changes were further explained by Yang et al. [33] who established that long-term DM induces oxidative stress and lipid peroxidation with impairment of membrane functions. This occurs through decreasing membrane fluidity and changing the activity of membrane-bound enzymes and receptors. Additionally, Li et al. [19] clarified neuronal cells’ apoptosis, as diabetic hyperglycaemia leads to opening of mitochondrial permeability transition pores which directs up expression of caspase-3 and caspase-8 that play a central role in the execution-phase of cell apoptosis.

Seeking for potential improving role of MSCs, fluorescent-labelled MSCs were recognised in the hippocampus after their systemic injection according to each DM + MSCs subgroups. In the current work, the histological and ultrastructural examination of MSCs treated subgroups showed variable amelioration of the pathological changes and increase in neuronal numbers regarding the extent of MSCs treatment and degree of damage produced by DM on the hippocampus. MSCs treatment for 4 weeks after 4 weeks of diabetic neuronal damage led to improvement of apoptotic changes which was more evident in CA1 region than in CA3 region. Moreover, this improvement was more apparent with prolonged duration of MSCs treatment to 8 weeks after 4 weeks of diabetic neuronal damage. There was obvious regression of neuronal apoptotic changes of CA1 and CA3 with relatively normal pyramidal neuronal structure.

Those histological changes were further documented by the statistical analysis of the morphometric counting of pyramidal neurons of both regions.

[27] Orlovsky et al.
[33] Yang et al.
[36] Zhao et al.
[31] Diabetes mellitus
[35] MSCs treatment
[19] Li et al.
There was increase in neuronal numbers especially with prolonged MSCs treatment in both regions, but this increase was still lower than in control groups. In the reviewed literature, no studies examined the effect of i.v. MSCs administration on hippocampal neuronal diabetic degeneration. However, the MSCs induced neuronal improvement of CA1 region after short-term treatment observed in the current work corresponds to findings reported by Kumar et al. [16] who studied cold stress as a cause of hippocampal neuronal degeneration. Also, the improvement found after longer duration of MSCs treatment was supported by the results obtained by Ye et al. [34] who studied the neuronal degeneration in Parkinson disease, as there was neuronal recovery after 10 weeks of MSCs transplantation. While, the moderate improvement observed in the current work in CA3 region was in harmony with the findings of Matchynski-Franks et al. [24] who studied MSCs effect after 10 weeks of transplantation in a mouse model of Alzheimer’s disease.

Discussing the improving role of MSCs in the current study to ameliorate diabetic neuronal degeneration and functional impairment, many authors explained their effects in the injured brain. Mezey et al. [25] concluded that human MSCs could enter the human and rat brains and generating neurons especially in the hippocampus and cerebral cortex. Moreover, Zhang et al. [35] found that transplanted MSCs could engraft into brain tissue and differentiate in vivo into neurons and glial cells. In contradiction, Sanchez-Ramos et al. [30] described that a small proportion of MSC-derived cells differentiated into neuron-like cells. Those findings were explained by Crigler et al. [7] and Munoz et al. [26] as MSCs implanted into the hippocampus did not proliferate, but they greatly increased proliferation, migration and differentiation of the endogenous neuronal stem cells after 8 weeks of neuronal degeneration. Prockop [28] described that MSCs could promote the neurogenesis of primary neural progenitors and survival of neural cells by expressing neurotrophic factors. Also, Calió et al. [4] reported that MSCs could decrease neuronal oxidative stress and apoptosis changes by enhancing the anti-apoptotic activity and antioxidant protection. These results were very promising for the development of a new therapeutic strategy for patient exhibiting diabetic neuronal degeneration and learning and memory impairment by using MSCs treatment.

CONCLUSIONS

Administration of MSCs ameliorates diabetic neuronal degeneration and learning and memory impairment mainly through promoting neurogenesis, alleviating oxidative stress and also through its anti-apoptotic effects. So, treatment with bone marrow-derived MSCs may exert a successful curative role in patients suffering from diabetic neuronal complications and improving their quality of life.

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REFERENCES


