Bone marrow stromal cells-derived exosomes target DAB2IP to induce microglial cell autophagy, a new strategy for neural stem cell transplantation in brain injury

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Abstract. Bone marrow stromal cells (MSCs) are a useful source of stem cells for the treatment of various brain injury diseases due to their abundant supply and fewer ethical problems compared with transplant treatment. However, the clinical application of MSCs is limited due to allograft rejection and immunosuppression in the process of MSCs transplantation. According to previous studies, microglial cell autophagy occurs following co-culture with MSCs. In the present study, exosomes were obtained from MSCs and subsequently characterized using transmission electron microscopy, atomic force microscopy and dynamic light scattering particle size analysis. The type of microRNAs (miRs) found in the exosomes was then analyzed via gene chip. The results demonstrated that microglial cell autophagy could be induced by exosomes. This mechanism was therefore investigated further via reverse transcription-quantitative PCR, western blotting and luciferase assays. These results demonstrated that exosomes from MSCs could induce microglial cell autophagy through the miR-32-mediated regulation of disabled homolog 2-interacting protein, thus providing a theoretical basis for the clinical application of miRs in MSCs.

Introduction

Conventional drug therapy cannot activate neurons in the brain (1). To treat neurological diseases such as encephalopathy, the only effective form of treatment is neural stem cell transplantation (1-3). Neural stem cells are undifferentiated pluripotent or committed cells (4), the transplantation of which has been documented as an effective method to repair and replace damaged brain tissue by reconstructing partial loop and function (5). Kochanek et al (6) transplanted neural stem cells into the damaged brain tissue of a Parkinson’s disease rat model and demonstrated that tremor symptoms were significantly mitigated, in a manner that may be associated with the production of dopamine in midbrain neural stem cells. In another study, Ogawa et al (7) cultured E14.5 stem cells derived from embryonic spinal cords in vitro, after which histopathological analysis was performed 5 days after transplantation. The results revealed that treated rats had significantly recovered motor function. However, immune reactions induced by neural stem cell application and the resulting rejection continue to be the main challenges obstructing their current clinical application (8,9). Therefore, it is important to understand the mechanism underlying the suppression of immune-related cell activity during neural stem cell transplantation (10,11).

Exosomes serve as important tools for the exchange of biologically active substances between cells (12). Most, if not all, mammalian stem cells release exosomes containing certain characteristics of their source parent cells (13). Exosomes contain cell-specific proteins, lipids and nucleic acids, which act as carriers for signal transduction, information exchange between cells and participate in the development of normal physiological processes and diseases (14). Li et al (15) previously reported that human umbilical cord mesenchymal stem cell exosomes significantly inhibit the ratio of peripheral blood CD3⁺CD4⁻T cells and CD3⁺CD8⁺T cells in normal humans. Exosomes with effective immunosuppressive functions have been demonstrated to provide a novel target for immunosuppression in transplantation settings, including bone marrow transplantation (16), cardiac transplantation (17), and liver transplantation (18). Moreover, exosomes are also involved in the immune responses against tumors (19) and have been shown to be effective in the treatment of chronic inflammatory diseases such as multiple sclerosis (20). In addition, exosomes derived from MSCs can inhibit the development of graft-versus-host disease (21,22). These studies suggest that exosomes may provide a new treatment option for the treatment of brain injury diseases.
for immunotherapy in treating tumors and autoimmune diseases (16,17).

Previous studies have suggested that autophagy participates in the regulation of inflammation to prevent the development of autoimmune and inflammatory diseases (18). Autophagy not only eliminates macromolecules in autophagic cells, but also clears damaged organelles to maintain intracellular homeostasis (19). Microglia are an important type of neuroimmune cell, which in their activated state, induce tissue repair and neuroprotection by releasing neurotrophic factors and phagocytizing damaged nerve cells (20). In cases of acute trauma to the central nervous system, including traumatic brain/spinal injury, hypoxia or ischemic brain damage, microglia rapidly initiate an immune response (21). Appropriate activation of microglia is beneficial for wound repair and microenvironmental reconstruction, which serves an important role in a number of nerve cell repair processes (22). The occurrence of autophagy in microglia also serves an important role in the differentiation, survival and homeostasis maintenance of transplanted stem cells (23). A study by Wang et al indicated that bone marrow-derived neural progenitor cells can differentiate into neurons, the transplantation of which in vivo can effectively promote motor function in rats following brain injury (24).

In previous studies, bone marrow-derived neural progenitor cells have been characterized, revealing that these cells have the potential to differentiate into neurons (25‑27). However, progress has been slow regarding investigation into the treatment of brain injury using neural stem cell transplantation, which may be due to changes in the intracranial microenvironment following brain injury (26). A series of studies have reported that the autophagy of microglia serves an important role in brain injury, involving cranial nerve inflammation, cerebral ischemia and cerebral hypoxia (28‑30). Stem cells that are transplanted into the body frequently fail and do not result in tissue repair (31). This may be due to the fact that stem cell transplantation is an exogenous procedure. Whether this process activates microglia autophagy, or whether microglia autophagy is associated with this process is yet to be fully elucidated. Observation and study on this series of problems are therefore urgently required for future clinical work on cell transplantation. To expand on previous studies assessing bone marrow‑derived neural progenitor cell-mediated tissue repair (28‑30,32), the present study systematically characterized the structure and size of bone marrow‑derived neural progenitor exosomes using optical technology, analyzed its content using second‑generation sequencing technology and investigated the molecular mechanism underlying microglia autophagy induced by the exosomes from bone marrow‑derived neural progenitor cells using molecular and cell biology techniques. The present study provided theoretical information on neural progenitor cell survival and differentiation following the transplantation of bone marrow‑derived neural progenitor cells, in addition to offering mechanistic and experimental support for the future clinical application of cell transplantation.

Materials and methods

Materials. All reagents and chemicals were purchased and used directly without further purification. The bone marrow stromal cell line was collected from the rat model of our team (28‑30), whilst the BV-2 microglial cell line was provided by CHI Scientific Inc. (cat. no. 7-1502). All aqueous solutions were prepared in deionized water and triple distilled water was used for all in vitro procedures. MTT, penicillin and trypsin were purchased from Sigma-Aldrich; Merck KGaA. DMEM/F12 and FBS were purchased from Thermo Fisher Scientific, Inc. and Zhejiang Tianhang Biological Technology Co., Ltd., respectively. ExoQuick™ reagent (cat. no. EXOQSA-1; Guangzhou Ruijing Information Technology Co., Ltd.), bicinechonic acid (BCA) protein assay kit (cat. no. P0012S; Beyotime Institute of Biotechnology) and Ultrafiltration centrifuge tubes (cat. no. UFC901096) were purchased from Guangzhou Ruijing Information Technology Co., Ltd. Rabbit antibodies for disabled homolog 2-interacting protein (DAB2IP; cat. no. ab87811), Beclin1 (cat. no. ab62557), microtubule-associated protein 1A/1B-light chain 3 (LC3; cat. no. ab48394), p62 (cat. no. ab109012), CD81 (cat. no. ab124717), CD9 (cat. no. ab223052) and β-actin (cat. no. ab79467) were purchased from Abcam. The present study was approved by the Ethics Committee of Guangdong Provincial People's Hospital (approval no. GDREC2018042A).

Isolation and culture of bone marrow‑derived neural progenitor cells. The methods and procedures for the isolation, culture and identification of bone marrow‑derived neural progenitor cells were performed based on previous studies (25‑27). Three Sprague-Dawley rats (sex, male; age, 5 weeks; weight, 115‑150 g) were housed in a temperature of 18‑25°C and 30‑60% humidity. Temperature 18‑25°C, humidity, 30‑60%. Animals were purchased from the Guangdong Medical Laboratory Animal Center. The rats were sacrificed by cervical dislocation and placed in a beaker containing 75% alcohol for 10 min. The bilateral femurs and tibias were cut, and the soft tissues were removed. The medullary cavities of the bones were washed with α-MEM supplemented with 10% fetal bovine serum. Centrifugation occurred at 200 x g for 20 min. The cloudy cell layers were plated on attachment plastic tissue culture dishes, and the α-MEM containing 10% FBS was changed 4 days later and renewed every 3 days. The cells were plated to obtain adherent rat bone marrow stromal cells (MSCs), which were harvested when cells reached 60‑80% confluence.

Extraction of exosomes from bone marrow‑derived neural progenitor cells. Neural progenitor cells were cultured in a 5% CO₂ incubator at 37°C. When the confluence reached 70‑80%, cells were washed with physiological saline and DMEM/F12 medium was replaced. The medium was changed after 48 h to collect the cell supernatant, followed by centrifugation (5,000 x g for 20 min, 4°C) to remove cells or cell debris. The exosomal supernatant (12 ml) was then added to the ultrafiltration centrifuge tube, and another 12 ml of the supernatant was added into another ultrafiltration centrifuge tube and centrifuged at 5,000 x g for 40 min (4°C). The supernatant was subsequently transferred to a 1.5 ml sterile, enzyme‑free centrifuge tube, which was covered well. ExoQuick Exosome reagent was subsequently added. The tube was turned upside down to ensure adequate mixing and incubated at 4°C for 12 h. Subsequently, the mixture was centrifuged at 1,500 x g for
30 min (4°C), before the supernatant was carefully removed and the remaining mixture was centrifuged (5,000 x g, 25°C) for another 5 min. All the liquid was carefully discarded, leaving only the white portion deposited at the bottom. Finally, 10% original volume of DEPC water was used to resuspend the pellet and mixed prior to storage at 4°C for further use.

**Identification of exosomes from bone marrow-derived neural progenitor cells and microRNA (miRNA) sequencing.** The morphology of exosomes was observed by transmission electron microscopy. A total of 10 µl of the 50 µl freshly extracted exosomal suspension was obtained for a hanging drop preparation on a copper mesh for 10 min and dried using filter paper. Subsequently, 5 µl of 2% phosphotungstic acid negative dye solution was added for 5 min and washed twice with ultrapure water. Excess water was absorbed with filter paper. The copper mesh was then incubated at room temperature until the sample dried naturally. Samples were then placed into sample boxes for observation under a transmission electron microscope at 20°C and 30% humidity (Hitachi H-7650, Hitachi, Ltd.). Exosomes were fixed with 90% ethanol at 20°C for 10 min. Data were analyzed using DigitalMicrograph 3.9 (Gatan, Inc.).

Exosome morphology was observed using atomic force microscopy (AFM). Freshly extracted exosomal suspension was first diluted with 0.9% physiological saline. A freshly cleaved mica plate was then fixed onto the surface of a glass slide, where 50 µl sample solution was dropped slowly onto the mica plate and allowed to dry. Finally, the sample was examined at room temperature using a silicon probe in tapping mode (Dimension Edge; Bruker Corporation).

The size and distribution of the freshly prepared exosomes (300 µl) were measured using a particle size analyzer (Zetasizer Nano ZS90; Malvern Instruments, Ltd.). The ID50 was obtained and calculated once the cumulative size distribution percentage reaches 50% of the grain size.

For on-column DNase digestion, exosome sample solution (400 µl) was added to 400 µl 2X RNAgents® Denaturing Solution (cat. no. Z5651; Promega Corporation), mixed by vortexing and incubated on ice. An equal volume of phenol/chloroform was added, mixed by vortexing and centrifuged (5,000 x g for 20 min) at 4°C. The supernatant was then pipetted into absolute ethanol (1 ml), mixed and transferred to a column for centrifugation (12,000 x g for 10 min) at 4°C. A total of 350 µl miRNA wash solution 1 was added, followed by further centrifugation (12,000 x g for 5 min at 4°C) and the supernatant was discarded. DNase I (10 µl) was mixed with 70 µl RDD buffer (Qiagen GmbH) before being transferred to the spin column and placed at room temperature for 15 min. A total of 350 µl miRNA wash solution 1 was added before further centrifugation (12,000 x g for 5 min at 25°C) to discard the supernatant. Wash solution 2/3 (500 µl) was then added to rinse the column twice. The spin column was then placed into a new collection tube, where 100 µl pre-warmed elution solution was added to the center of the column, followed by centrifugation (12,000 x g for 5 min at 25°C). The resulting liquid in the collection tube was the extracted total RNA and the sample was stored at -80°C.

Total RNA was initially used in a chip experiment, which was subjected to quality inspection using NanoDrop ND-2000 spectrophotometer (Thermo Fisher Scientific, Inc.) and Agilent Bioanalyzer 2100. Qualified RNA was then subjected to subsequent chip experiments. High-throughput analysis of the miRNA expression profiles was performed using Agilent miRNA chip technology (Rat miRNA 8x15k v2.1 microarray; Agilent Technologies GmbH). Sinotech Genomics performed miRNA sequencing. The chip results were scanned using the Agilent Microarray Scanner, where the data were read by the Feature Extraction software 10.7.1.1 (Agilent Technologies GmbH), followed by normalization using the R language package.

**Bone marrow-derived neural progenitor exosome-induced microglia autophagy.** For autophagy detection in BV2 cells, different amounts of exosomes were added to BV2 cells (5 µl, 360 ng/µl; 15 µl, 360 ng/µl) at 37°C. After 48 h, an inverted fluorescence microscope (TE2000-E; Nikon Corporation) was used to capture images. The culture medium (DMEM/F12) was then removed, cells were washed once with PBS and fixed with 2.5% glutaraldehyde solution at 4°C overnight. The fixative was removed, the sample was rinsed three times with PBS and fixed with 1% osmic acid solution for 2 h at 37°C. The sample was subsequently dehydrated using an ascending ethanol gradient (30, 50, 70, 80, 90 and 95%) and treated with pure acetone for 20 min at 25°C. The sample was treated with a mixture of epikote resin and acetone (v/v=1/1) for 1 h, followed by the same mixture but at a higher concentration of the embedding agent (v/v=3/1) for 3 h before incubation with 100% embedding agent overnight at 25°C. The treated sample was embedded and heated at 70°C overnight to obtain an embedded sample. The samples were sectioned using a Leica EM UC7 ultramicrotome (Leica Microsystems GmbH) where 70-90 nm sections were obtained. The sections were stained with 10% lead citrate solution for 10 min at 25°C and 50% uranyl acetate for 10 min at 25°C before observation under a transmission electron microscope (Hitachi H-7650, Hitachi, Ltd.).

For colony formation assays, BV cells were seeded into six-well plates at 500 cells/well. Following 24 h incubation at 37°C with 5% CO₂, DMEM was then replaced and exosomes were added for 14 days (5 µl, 360 ng/µl). The control group was not treated with exosomes. The cells were then fixed with 90% ethanol for 30 min at 25°C and washed with PBS three times. After further incubation for 30 min at 37°C in the dark and staining with 1X Giemsa solution at 25°C for 20 min, images was obtained on an inverted microscope (TE2000-E; Nikon Corporation) at x100 magnification.

**MTT assay.** Inhibition of cell growth by exosomes was measured using an MTT assay. The cells were exposed to exosomes at different times (12, 24 and 48 h). Briefly, BV cells were seeded in 96-well tissue culture plates (5x10³ cells/per well) for 12 h at 37°C with 5% CO₂. After exosome (5 µl, 360 ng/µl) treatment, BV2 cells were incubated for 24 h at 37°C with 5% CO₂. After incubation, 20 µl/well MTT solution (5.0 mg/ml in PBS) was added and incubated for 4 h at 37°C with 5% CO₂. The medium was aspirated and replaced with 100 µl/well DMSO to dissolve the formazan salt. The absorbance intensity was measured by using a microplate spectrophotometer at a wavelength of 570 nm (Thermo Fisher Scientific, Inc.).
Molecular analysis of autophagy induction. The number of exosomes used for molecular mechanism analysis was determined by measuring the total protein concentration in the exosomes, which was obtained using the BCA kit (360 ng/µl).

Western blotting. The exosomes secreted by the bone marrow-derived neural progenitor cells and the bone marrow-derived neural progenitor cells of the normal control group were first collected by centrifugation at 12,000 x g for 10 min at 4°C. A total of 70 µl RIPA buffer (cat. no. P0013C; Beyotime Institute of Biotechnology) was then added and incubated on ice for 30 min. The protein was quantified using the BCA method, following which the protein mass concentration was adjusted to 5 g/l using RIPA buffer and stored at -80°C for further use. For electrophoresis, 20 µg protein was separated by 10% SDS-PAGE, before being transferred onto nitrocellulose membranes at 100V for 1 h and blocked in blocking solution (cat. no. P0216; Beyotime Institute of Biotechnology) for 1 h at 37°C. The membranes were then incubated with an appropriate amount of primary antibodies against DAB2IP, Beclin1, p62, CD81, CD9 and LC3 (all, 1:1,000) overnight, following which the membranes were incubated with corresponding secondary antibody (alkaline phosphatase-labeled goat anti-rabbit immunoglobulin G; 1:1,000; cat. no. A0239; Beyotime Institute of Biotechnology) for 1 h at room temperature. The protein bands were visualized by ECL luminescence, following which densitometric analysis was performed. Protein bands were visualized using GelDoc XR (BioRad Laboratories, Inc.).

Calcium ion detection. Different amounts of exosomes were added to BV2 cells (5 µl, 360 ng/µl; 15 µl, 360 ng/µl), which were then incubated for 24 h at 37°C. Fluo-3 AM working solution (1 µM; cat. no. S1056; Beyotime Institute of Biotechnology) was added to sufficiently cover the cells. After incubation for 30 min at 37°C, the fluorescence of Fluo-3 was measured (excitation wavelength, 488 nm; emission wavelength, 530 nm) using a fluorescence microscope plate reader (Varioskan Flash; ThermoFisher Scientific, Inc.) to assess any changes in intracellular calcium ion concentrations.

Luciferase assay. The miRNA and 3'-UTR binding site (accession no. NM_032552.3; Homo sapiens DAB2 interacting protein, DAB2IP, transcript variant 1, mRNA) was predicted by BLAST (https://docs.microsoft.com/en-us/previous-versions/hh397746(v=msdn.10)?redirectedfrom=MSDN). Upon reaching a confluence of ~70%, 293T cells (American Type Culture Collection; incubated at 37°C with 5% CO2) were seeded into six-well plates at a density of 2x10^5 cells/well, where 2 ml Opti-MEM was added to each well, and the fluorescence of Fluo-3 was measured for a second time. Renilla luciferase activity was used as an internal control.

Reverse transcription-quantitative PCR (RT-qPCR). BV2 cells were seeded in six-well plates (1x10^5 cells/well) for 12 h at 37°C, then different amounts of exosomes were added to BV2 cells (5/15 µl, 360 ng/µl) for 24 h at 37°C. The BV2 cells were collected, total RNA was then extracted using RNAprep Pure Cell/Bacteria kit (cat. no. DP430; Tiangen Biotech Co., Ltd.) in accordance with the manufacturer's protocol, following which RNA purity was measured using a NanoDrop 2000 spectrophotometer (Thermo Fisher Scientific, Inc.). The sequences used were as follows: U6 forward, 5'-CTCGCTTCGGCAGCAC-3' and reverse, 5'-AACGCTTCACAGAAAACTTGAGTTGTTGATAT-3'; miR-32-3p, 5'-CAATTTGATGTTGATATT-3'; inhibitor-miR-32-3p, 5'-AAATATCAGACAAGTACATTTGAA-3'; miR-188-5p, 3'-CATCCCCTTGATGGTGGGGG-5'; miR-211-3p, 3'-GCAAGGACAGAACCAAGGGGG-5'; miR-188-5p, 3'-CATCCCCTTGATGGTGGGGG-5'; miR-466b-5p, 3'-TGATGTTGTTGACATGACAT-5'.

RNA was added with polyA tail and reverse transcribed into cDNA (RevertAid First Strand cDNA synthesis kit; cat. no. K1622; Thermo Fisher Scientific, Inc.), following which the expression of four potential target genes were detected via qPCR (GoTaq qPCR and RT-qPCR Systems; cat. no. A6001; Promega Corporation). Reactions were performed on a fluorescence qPCR instrument (CFX Connect; cat. no. 1855200; BioRad Laboratories, Inc.). The following thermocycling conditions were used for the qPCR: Initial denaturation at 95°C for 10 min, followed by 50 cycles of 95°C for 15 sec and 60°C for 1 min. The relative mRNA levels of target samples to control samples were calculated according to 2^(-ΔΔCt) method (33).

Statistical analysis. Data are expressed as the mean ± standard error of the mean based on at least three independent experiments. Statistical analysis was performed using one-way ANOVA, followed by a Bonferroni post hoc test for multiple group comparisons or a Student's unpaired t-test for pairwise comparisons. P<0.05 was considered to indicate a statistically significant difference unless otherwise indicated. All analyses were performed using GraphPad Prism Software version 6.0 (GraphPad Software, Inc.).

Results

Isolation of bone marrow-derived neural progenitor cells and the effect in BV2 cells. The methods and procedures for the
isolation, culture and identification of bone marrow-derived neural progenitor cells were performed based on previous studies (25-27). As shown in Fig. 1A-a, the morphology of the isolated and identified bone marrow-derived neural progenitor cells appeared to be slightly. The mouse microglial cell line BV-2 was purchased and sub-cultured, which were in fusiform states (Fig. 1A-b). Exosome-associated target proteins, such as CD9 and CD81, are types of biomarkers for exosomes (34). As shown in Fig. 1B, the exosomes of bone marrow-derived neural progenitor cells were first characterized via western blotting, by examining the expression of exosome-associated target proteins CD9 and CD81 (13-15).

**Morphology and particle size distribution of exosomes from bone marrow-derived neural progenitor cells.** The exosomes of bone marrow-derived neural progenitor cells were collected via ultrafiltration concentration centrifugation and observed by transmission electron microscopy (Fig. 1Ca). They were found to be either circular or elliptical in shape, of homogeneous distribution, with double-layered lipid vesicle (arrows). The vesicles secreted by bone marrow-derived neural progenitor cells have clear vesicle membrane structures with higher internal dioptries compared with the background, consistent with the exosome features observed (34). Based on this, the size of exosomes from bone marrow-derived neural progenitor
Sequencing miRNA found in exosomes from bone marrow-derived neural progenitor cells. The exosomes of bone marrow-derived neural progenitor cells were collected and total exosomal RNA was extracted. High-throughput analysis of miRNA expression profiles was performed using Agilent miRNA chip technology. An increasing number of studies have previously demonstrated that miRNAs serve important roles in various diseases and biological functions (34,35). miRNAs can regulate a number of physiological processes, including proliferation, cell cycle progression, apoptosis, invasion and angiogenesis by inhibiting the expression of target genes (36). The results of sequencing of miRNA shown that 758 miRNAs found in the exosomes met the quality control requirements of bone marrow-derived neural progenitor cells. The original data of chip-based detection is presented in Fig. 2B. The obtained data is demonstrated in Fig. 2C. The signal value of the original data miRNA probe is presented in Fig. 2D. From the 758 miRNAs detected in the exosomes of bone marrow-derived neural progenitor cells, the top 20 miRNAs according to abundance were selected. Information, including type name, the degree of correspondence between the probe

Figure 2. Sequencing miRNA isolated from exosomes from bone marrow-derived nerve progenitor cells. (A) Quality inspection result of total RNA isolated from the exosomes. (B) The original data obtained from the Agilent miRNA chip. Green dots show miRNAs that were detected. (C) The data of the Agilent miRNA chip normalized to the fluorescent signal point of the chip probe. (D) The fluorescence detection signal of the miRNA chip. miRNA, microRNA. BG, background.
and the original data, in addition to their active sequence structure information, are provided in Table I. After combining literature reports (37-41), four potential miRNAs which have been documented to serve a role in exosome-induced microglia autophagy were identified: rno- miR-32-3p, rno- miR-211-3p, rno- miR-188-5p and rno- miR-465-5p. RT-qPCR was performed to investigate the differences in the relative expression of these four potential target miRNAs in the microglia cells in the presence or absence of bone marrow-derived neural progenitor exosomes (Fig. 3).

It was revealed that miR-32-3p exhibited the highest expression, which meant it could be used as a potential target molecule (Fig. 3A). Therefore, the expression of miR-32-3p in microglia was examined further using different volumes of exosomes and different treatment times, the results of which are presented in Fig. 3B.

**Bone marrow-derived neural progenitor exosomes induce microglia autophagy.** Using molecular biology and cell biology techniques, the mechanism underlying microglia autophagy induced by exosomes from bone marrow-derived neural progenitor cells was investigated further. MTT and colony formation assays were performed to investigate the effects of exosomes from bone marrow-derived neural progenitor cells on the viability and proliferation of microglia. Exosomes of bone marrow-derived neural progenitor cells were collected during microglia growth (Fig. 4A). High-dose exosomes (Ex-H) had no effect on the growth of microglia. No statistically significant differences were observed in the survival rates between cells cultured for 12 h with exosomes and cells without the addition of exosomes, nor between cells cultured in the absence of exosome treatment and those treated with exosomes for 24 and 48 h.

This suggested that exosomes of bone marrow-derived neural progenitor cells had no significant effect on microglia cell growth. This can be clinically relevant if performed without affecting the growth of normal cells. By measuring the formation rate of colonies, the effect of exosomes from bone marrow-derived neural progenitor cells on the proliferation of microglia was investigated (Fig. 4B). The number of microglial

| Systematic name | Total probe signal (raw) data | Total probe signal (normalized) | Sequence |
|-----------------|-------------------------------|---------------------------------|----------|
| miR-1224        | 1511.147                      | 10.561428                       | 5'-GTGAGGACTCGGGAGGTGG-3' |
| miR-466b-5p     | 737.30896                     | 9.526125                        | 5'-TATGTGTGTGTATGTCCATG-3' |
| miR-32-3p       | 129.2438                      | 7.013951                        | 5'-CAAATTTAGTGTGTGTGATAAT-3' |
| miR-1896        | 82.5686                       | 6.3675213                       | 5'-TGGGGGTTGAGGAGGAGG-3' |
| miR-672-5p      | 73.4067                       | 6.1978397                       | 5'-TGGAGGTTGGTGTGACTGTGGTGA-3' |
| miR-1306-3p     | 64.6163                       | 6.013826                        | 5'-ACGTTGGCTCTGGTGGT-3' |
| miR-211-3p      | 34.9084                       | 5.128681                        | 5'-GCGAGGACAGGAAAGGGTGTC-3' |
| miR-466c-5p     | 33.4107                       | 5.062238                        | 5'-TGTGATGTTGTATGTACATG-3' |
| miR-466d        | 31.84107                      | 4.992817                        | 5'-ATGTTGTTGTATGTCTTTT-3' |
| miR-465-5p      | 31.7184                       | 4.987248                        | 5'-TATTTAGAAGGTTGCTGTGTTG-3' |
| miR-3473        | 31.4372                       | 4.9744005                      | 5'-TCTAGGGCTTGAGAGGCTCA-3' |
| miR-3593-3p     | 27.8128                       | 4.797677                        | 5'-CCATCACACACAGACTGGATTTTG GCCCGAGGGTTAGCTGCTAGAGAGAGAGGAGTGG-3' |
| miR-1249        | 26.360802                     | 4.720322                        | 5'-GGGAGGAGGAGGAGGAGATGGGCCAAGT TCCCTCTGGCTGGAAGCCTCCCCCCTCTTCACCTG-3' |
| miR-6216        | 25.43394                      | 4.668683                        | 5'-GATACACAGGGCAGGAGGAGAA-3' |
| miR-3588        | 25.2884                       | 4.660381                        | 5'-TCACACAGGTAGCTGCAAGG-3' |
| miR-483-5p      | 24.348429                     | 4.6057568                       | 5'-AAGAGGGGAGAAGAGAGAGGAG-3' |
| miR-2985        | 23.6542                       | 4.5640244                      | 5'-ATGCCACACCTAAAGGATCCCTATTA AGGTGGTGGGTTGATAGTATAACAATGTGCT CAAATGTTGTATATAGTATACCCACCTACCTG ATGTTGCTTTTAAGACTCTAACGG-3' |
| miR-296-5p      | 20.11955                      | 4.3305264                      | 5'-AGGGCCCCCCCCCTAATCCTTG-3' |
| miR-188-5p      | 19.535389                     | 4.2880177                      | 5'-CATCCCCCTTGATGGTGAGGG-3' |
| miR-3562        | 18.44011                      | 4.2047753                      | 5'-TTGGGGCGACTGGTGGATGGGA-3' |

miR, microRNA.
colonies formed following the addition of exosomes from bone marrow-derived neural progenitor cells was not found to be markedly different from the control group, suggesting that the addition of exosomes from bone marrow-derived neural progenitor cells did not affect microglial proliferation.

Notably, the shape of microglial cells change from roundness to fusiform after the addition of exosomes from bone marrow-derived neural progenitors (Fig. 4C). Damaged organelles such as swollen and degenerated mitochondria were observed under a transmission electron microscope (Fig. 4D). Additionally, residue that could not be fully degraded in the autolysosome was observed. These results suggested that exosomes from bone marrow-derived neural progenitor cells induced microglia autophagy without affecting growth and proliferation.

Western blotting and luciferase reporter gene assays were subsequently performed to validate the potential targets of miR-32-3p and associated regulatory mechanism. Following the addition of exosomes from bone marrow-derived neural progenitor cells, the expression of the DAB2IP protein in microglial cells was found to be upregulated whilst that of p62 was significantly downregulated (Fig. 5A). The shear band (the upper or lower band LC3 II/LC3 I) of the LC3 protein was also found to be significantly increased. In addition, Beclin 1 was observed to be significantly upregulated following the addition of exosomes from bone marrow-derived neural progenitor cells (Fig. 5A). Different amounts of exosomes were added to BV2 cells [Low dose (Ex-L), 5 µl, 360 ng/µl; high-dose (Ex-H), 15 µl, 360 ng/µl], and the expression of proteins CD9 and CD81 of Ex-H was markedly higher compared with Ex-L (Fig. 1B). These results suggested that microglia autophagy may be associated with the target binding of miR-32-3p and the DAR2IP (39), alongside the subsequent activation of Beclin1 (42).
The potential process of endoplasmic reticulum stress was examined further. As presented in Fig. 5B, after treatment with exosomes, the intracellular Ca\textsuperscript{2+} level in BV2 cells significantly increased compared with controls. The potential interaction between miR-32-3p and the DAB2IP protein was examined further using luciferase reporter gene assays. As shown in Fig. 5C, miR-32 could bind DAB2IP by directly targeting 3'-UTR, to regulate the expression of DAB2IP mRNA.

Discussion

A number of studies have demonstrated the important role of microglia autophagy in brain injury, including cranial nerve inflammation, cerebral ischemia and cerebral hypoxia (28-30). Clinical stem cells transplantation frequently fails and does not result in tissue repair (31). It is therefore important to investigate the effect of microglia autophagy in this process. Observation and study on this series of problems will be helpful in exploring future clinical applications regarding cell transplantation. Microglia autophagy serve an important role in the differentiation, survival and homeostasis maintenance of transplanted stem cells (43-46). Previous studies have demonstrated that bone marrow-derived neural progenitor cells can differentiate into neurons and their transplantation \textit{in vivo} can effectively promote the motor function of rats following brain injury (25-27).

Microglial cell autophagy has also been found in the process of bone marrow-derived neural progenitor cell transplantation during the treatment of brain nerve injury in SD mice in another previous study (47). Investigating the relationship and mechanism between microglia autophagy and the survival and differentiation of neural stem cells are of significance for targeting microglia-associated inflammation, promoting the survival of transplanted neural stem cells in damaged tissues and improving their ability to repair tissues.

The number of studies into the role of exosomes on biological processes is increasing where their importance is gradually being revealed (48,49). However, since the extraction yield of exosomes reported in existing studies has generally been low, the specific molecular mechanism of exosome involvement in intercellular communication remains unclear (50).

Therefore, using information obtained from previous studies, the present study extracted exosomes of high-yield (51,52) and high purity from bone marrow-derived neural progenitor cells. The molecular mechanism of the effects of autophagy on the relationship between bone marrow-derived neural progenitor cells and microglia was subsequently analyzed by sequencing and western blotting. The pathway of microglia...
autophagy that was affected by exosome miRNA was then verified, where a potential molecular mechanism of action was proposed. Results of present study provided a basic theoretical basis for an in-depth study on the clinical application of RNA molecules in exosomes secreted by bone marrow-derived neural progenitor cells.

Among the 758 miRNAs detected in the exosomes of bone marrow-derived neural progenitor cells, the top 20 miRNAs were selected according to detection abundance. Their type name, the extent of probe correspondence between original and normalized data, and the active sequence structure information were obtained. Combining bioinformatics analysis and literature reports, four potential miRNAs (rno-miR-32-3p, rno-miR-211-3p, rno-miR-188-5p and rno-miR-465-5p), which have been reported to serve a role in exosome-induced microglia autophagy, were identified (53,54). Using RT-qPCR, differences in the relative expression of these four potential target miRNAs in microglial cells in the presence and absence of exosomes of bone marrow-derived neural progenitor cells was investigated further. Following the addition of exosomes from bone marrow-derived neural progenitor cells, all four potential target miRNAs exhibited significantly increased expressions in microglial cells, where the expression of miR-32-3p, which is located in the 14th intron of the gene C9orf5, was found to be the highest, suggesting that it is a potential target molecule. Previous studies on the biological effects of miR-32 have mainly concentrated on tumors, viral replication and androgen regulation, which demonstrated that upon miR-32 overexpression, growth of the androgen-dependent prostate cancer cell line LNCaP was promoted. miR-32 has also been found to reduce apoptosis by directly regulating the expression of the $\text{BTG2}$ gene (55). Upregulation of miR-32 has been reported to inhibit the expression of apoptosis-inducing protein phosphoinositide-3-kinase interacting protein 1 and regulate the Ser/Thr protein kinase mitogen-activated protein kinase kinase, thus serving an important role in a variety of different signal transduction pathways (56). miR-32 serves a regulatory role in a number of physiological processes, including cell growth, differentiation, inflammation, apoptosis, vascular endothelial cell proliferation and neovascularization (57,58). However, the role of miR-32 in microglial growth, proliferation and apoptosis remains poorly understood.

In the present study, molecular and cell biology techniques revealed that exosomes from bone marrow-derived neural
progenitor cells exerted no significant effects on the growth and proliferation of microglia. Notably, following the addition of exosomes from bone marrow-derived neural progenitor cells, microglia began to develop autophagy, with typical autophagy features (59) observed under light and transmission electron microscopy. The results suggested that exosomes from bone marrow-derived neural progenitor cells can induce microglia autophagy without affecting the growth and proliferation of microglia.

Western blot and luciferase reporter gene assays were performed to validate the potential targets of the miR-32-3p associated regulatory mechanism. Following the addition of exosomes from bone marrow-derived neural progenitor cells, DAB2IP protein expression in microglia was found to be upregulated. The expression of Beclin1, which has been recently determined to be involved in the regulation of autophagy (60), was also found to be upregulated.

These results suggested that autophagy in microglia may be associated with miR-32-3p. In this process, miR-32-3p may be produced by the introduction of exosomes from bone marrow-derived neural progenitor cells. During the induction of autophagy in microglia, the marker protein p62 was degraded in the current study, whilst that of the LC3 protein shear band was increased, suggesting that the conditions for cellular stress to cause the endoplasmic reticulum to initiate autophagy.

The present study also examined the underlying process of endoplasmic reticulum stress. Following the addition of exosomes from bone marrow-derived neural progenitor cells, microglial intracellular calcium concentrations were increased to slightly increased. This suggested that the entry of exogenous signals resulted in the induction of endoplasmic reticulum stress in microglial cells, consistent with the degradation of p62 found in western blotting. Combining these experimental results, the mechanism underlying microglia autophagy induced by the exosomes from bone marrow-derived neural progenitor cells was hypothesized. First, miR-32-3p present in the exosomes secreted by bone marrow-derived neural progenitor cells enter into microglia, where they specifically bind to the DAB2IP to affect its expression. Then, the increase in DAB2IP protein expression in turn promotes the expression of the autophagy-inducing protein Beclin1 and enhances the stress response in microglia. At the same time, the endoplasmic reticulum initiates the stress response mechanism, increasing the intracellular calcium concentration (43) and causing p62 degradation, LC3 cleavage and the formation of cell autophagosomes. In this process, miR-32-3p was found to be a potential target molecule, whilst DAB2IP protein was a potential target protein. Briefly, bone marrow stromal cells (MSCs) are potentially useful for the treatment of a number of brain injury diseases due to their abundant supply. An increasing number of studies have reported their prospective functions and applications (9,12-17). In the present study, 758 miRNAs were detected in exosomes of MSCs using Agilent miRNA chip technology, where rno-miR-32-3p, rno-miR-211-3p, rno-miR-188-5p and rno-miR-465-5p were first found in the exosomes of MSCs.

Since microglia autophagy serves an important role in conditions associated with brain injury, including cranial nerve inflammation, cerebral ischemia and cerebral hypoxia, the results of the present study provided a theoretical reference for studies into brain injury treatment based on neural stem cell transplantation. In addition, the present study provided certain theoretical basis for cell survival and differentiation following transplantation of bone marrow-derived neural progenitor cells, in addition to theoretical and experimental support for future clinical cell transplantation. However, there are some limitations to the present study. For example, it only focused on the biological effects of exosomes from bone marrow-derived neural progenitor cells on microglia, instead of the effects of microglial metabolism. In addition, among the plethora of miRNAs found in the exosome, only 4 potential molecules were studied by bioinformatics analysis and the role of the remaining miRNAs on microglial autophagy cannot be ruled out. The present study only investigated the binding of miR-32-3p to DAB2IP instead of investigating its potential role further, which are the key issues that need to be addressed in the future.

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Availability of data and materials

The datasets used and/or analyzed during the current study are available from the corresponding author on reasonable request.

Authors’ contributions

FYY, MXZ, MSZ and JYO designed and directed the experiments. FYY, WFB and MSZ wrote the manuscript. All authors read and approved the final manuscript.

Ethics approval and consent to participate

The present study was approved by the Ethics Committee of Guangdong Provincial People’s Hospital (Certificate no. GDR EC2018042A).

Patient consent for publication

Not applicable.

Competing interests

The authors declare that they have no competing interests.

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