ABSTRACT

AIM: Glioblastoma multiforme (GBM) is one of the malignant brain tumors that occur most frequently. Despite advances in therapy techniques, the cure of GBM is a major concern. Accordingly, there is a lot of interest in devising novel approaches, such as stem cell therapy, to treat patients with GBM. The aim of this study was to investigate the effects of human bone marrow stem (BMS) cells as well as human olfactory ensheathing cells (OECs) on the outgrowth of U87 glioma in rats.

MATERIAL and METHODS: OECs and BMS cells were obtained from volunteers. After verification of the stem cell type by flow cytometry and immunocytochemistry (ICC), cells were labeled and injected into human glioma-bearing rats. Magnetic resonance imaging (MRI), Hematoxylin and Eosin (H&E), and Immunohistochemistry (IHC) were utilized to assess the properties of the groups.

RESULTS: We found extensive migration and homing of the OECs and BMS cells towards the tumor area. H&E and IHC staining indicated that the grafted OECs survived and prevented the development of glioma. BMS cells supported proliferation and new vessel formation, and metastasis in glioma tissue.

CONCLUSION: OECs and BMS cells can pass the blood brain barrier and reach the glioma mass. Therefore, this approach can be a potentially powerful method for the delivery of therapeutic agents to malignant brain tumors. In addition, these cells may be genetically modified in order to specifically express tumor-suppressive factors.

KEYWORDS: Glioblastoma multiforme, Stem cell therapy, Histopathology
OECs have high capacity of migration towards the damaged tissues of central nervous system (CNS) without tumorigenicity behavior (14,18). Therefore, the aim was to investigate the effects of bone marrow stem (BMS) cells as well as olfactory ensheathing cells on the outgrowth of U87 glioma in rats.

### MATERIAL and METHODS

This study was approved by the Ethical Commission of the Kerman University of Medical Sciences with Approval number EC/KNRC/86/23.

#### Culture of stem cells

**BMSCs**

BMS cells were extracted by processing bone marrow aspirates from one healthy donor as described previously (4). Briefly, the bone marrow was aspirated from the posterior iliac crest. The sample was rinsed, centrifuged, and Buffy coat was cultured at 37°C and 5% CO₂ air atmosphere. 48 hours later, the cells were washed; culture medium was added and then changed every 3 days. After 15 days, adherent cells formed colonies and when the cells achieved suitable confluency, the harvested cells from 3 passages were used in the different studies.

**OECs**

During a pituitary tumor resection from a 34-year-old male, the olfactory bulb was obtained (computed tomography (CT) and magnetic resonance imaging (MRI) findings of the patient showed a huge suprasellar lesion and the neurosurgeon had to cut the right olfactory bulb for better and more accurate surgery of the tumor). After isolating the olfactory bulb, it was stored in a falcon containing sterile Hank’s Balanced Salt Solution (HBSS) and antibiotic. The pia was removed and the olfactory bulb tissue was cut into small pieces. They were washed with HBSS and incubated for one hour at 37°C in a mixture of trypsin/DNase in HBSS. The tissue fragments were dissociated mechanically. Afterward, the dissociated cells were washed with phosphate buffered saline (PBS) and seeded onto culture flasks with complete medium. After 5 days, the flask was washed for removal of non-adherent cells, and culture medium was added. The harvested cells were detached with a mixture of trypsin/ Ethylenediaminetetraacetic acid (EDTA) and used in the study process.

#### Characterization of cells

**Flow cytometry of BMSCs**

The procedure surface of marker analysis was obtained from Seyed et al. (37). Briefly, the CD34, CD44, CD45, CD73, CD90, CD105, CD117, and OC4 markers were analyzed using fluorescence-activated cell sorting (FACS) in the BD Bioscience Flow cytometer.

**Osteogenic and Adipogenic differentiation of BMSCs**

Osteogenic and adipogenic differentiation methods were described elsewhere (21). Briefly, 2-3×10⁴ cells/cm² were seeded and fed with osteogenic and adipogenic inducing medium for 21 days. After 21 days, the Alizarin red and Oil-red staining process was performed in order to detect the Calcium-phosphate precipitates and fat droplets respectively.

#### Immunocytochemistry (ICC) of OECs

Immunocytochemistry procedure of the OECs was obtained from Salehinejad et al. (36). Briefly, cultured cells were fixated, washed, incubated with primary antibodies, including mouse monoclonal anti-S₁₀₀β (Chemicon) and mouse anti-p75 (Abcam, USA). The slides were washed and incubated with fluorescent secondary antibodies Alexa 594 and Alexa 488. Afterwards, slides were stained with Hoechst and mounted.

**Cells labeling with PKH₆₇**

PKH (a membrane- binding fluorescent dye) was employed to label bone marrow stem cells (BMSCs) and OECs. The procedure for labeling was described previously (1). Briefly, the cells were detached, centrifuged, and counted. Afterwards, cells were rinsed and suspended in Diluent C. Immediately, the staining mixture (Diluent C+PKH₆₇) was added to the cells. After 3-5 minutes, 400 µl fetal bovine serum (FBS) was added to stop the reaction. The cells were centrifuged, washed, and seeded in flasks.

**Culture and labeling of U87 Cells**

U87 MG cells were cultured at 37°C and 5% CO₂. Afterwards, cells achieved 80-90% confluency and were detached by trypsin/EDTA. U87 cells were centrifuged, counted, and used for the experimental procedure. To use the U87 cells in the experimental procedure, they were labeled with PKH₂₆ dye and the aforementioned labeling process (1).

**Induction of tumor**

In this study, 28 male Sprague-Dawley rats weighing 200-250 g were chosen. They were kept under normal conditions. They were randomly allocated into four groups (PBS, Tumor, BMSC, and OEC) of 7 rats. The implantation procedure of U87-PKH₂₆ cells was described previously (1). Briefly, rats were anesthetized and fixed in a stereotaxic frame. The scalp was incised and a hole was drilled in the skull. Then, the needle was inserted into the right frontal lobe. For three of the groups (except the PBS group), 10 µl of cell suspension (10⁶ cells) was injected. The same process was done for PBS group, but only 10 µl PBS was infused.

**Determination of tumor by MRI**

Ten days after tumor induction, MRI was performed on the rats as described previously (1). Briefly, the animals were anesthetized and positioned head-up in the coil of a 1.5-Tesla machine and both T1- and T2-weighted spin-echo images were obtained.

**Transplantation of PKH₆₇ cells**

The PKH₆₇ cells implantation process was described previously (1). Briefly, on the 10th day of the experiment, rats were anesthetized; the skin and muscles of the thorax were incised in the left fifth intercostal space. Afterwards, 2×10⁶ PKH₆₇ cells (100 µl) were infused by an insulin syringe into the left ventricle of the heart of the therapy group rats (BMSC and OEC).
Hematoxylin-Eosin (H&E) and Immunohistochemical (IHC) staining

On the 17th day of the experiment, all animals were anesthetized and perfused as described previously (1). The brains were post-fixed with the fixing/embedding, sectioning, and mounting process. Following deparaffinization/rehydration, some of the slides were stained with H&E and some of sections were used for the IHC staining process. For IHC staining, they were incubated with primary antibodies, including PECAM-1 monoclonal mouse anti-rat CD31, monoclonal mouse anti-human Nestin, monoclonal mouse anti-human Ki-67, and monoclonal mouse anti-human antibody glial fibrillary acidic protein (GFAP). Then, the slides were incubated with secondary antibodies; streptavidin/biotinylated peroxidase complex counterstained and mounted. The proportion of CD-31, Nestin, Ki-67, and GFAP positive cells, and the percentage of stained cells in 200 cells, were calculated.

For detecting PKH-cells in the animals, three brains from each tumor-bearing group were immediately frozen. Cryostat-generated slides were mounted and examined with a fluorescence microscope equipped with a digital camera.

Calculation of tumor volume

For this purpose, serial sections of each sample were prepared. The sections were 3 µm and the interval between the sections was 20 µm throughout the length of the tumor. These stained sections (with H&E) were photographed using a light microscope equipped with a digital camera. Serial images of each sample were used for calculating the tumor area (A) using the Analysis software. The Analysis software allows the reader to draw a line around the perimeter of the tumor. Each of these images, the tumor sectional areas were measured and the volume (V) was calculated by the following formula:

\[ V = \frac{1}{2} \sum (A_1 + A_2 + A_3 + \ldots + A_n) \times 20n + 3n. \]

Here, “n” is the number of sections containing tumor.

Statistical Analysis

Data are expressed as mean±SEM and the Kolmogorov-Smirnov test was used for measurement distribution of the data. Data were analyzed with the one-way ANOVA test (SPSS version 21) and a p value < 0.05 was considered significant.

RESULTS

Characterization of cells

BMSCs

Cultured BMS cells contained an analogous population of spindle-shaped cells (Figure 1A). Flow cytometry analysis revealed the expression of MSCs defined markers such as CD44 (54%), CD73 (82%), CD105 (80%), CD90 (83.33%), and negative expression of hematopoietic-specific markers CD34 (4.5%), CD117 (12.75%) and CD45 (5%). It has been shown that BMS cells express OCT4 (19%) (Figure 1B).

Under osteogenic induction medium, Alizarin Red staining revealed an extension of mineralization with detectable bone nodules in cells (Figure 1C). Interestingly, when BMS cells grew under adipogenic conditions, they showed fat droplets after staining by using oil-red O dye (Figure 1D).

OECs

In this study, OECs were successfully derived from the olfactory bulb, and the majority of OECs were spindle-shaped with a large nucleus. IHC data showed strong staining for Sβ and p75NGFR (Figure 2A-G).

MRI

MRI analysis revealed that a tumor mass was formed in glioma-bearing rats compared with control rats. On coronal T1-W images, the spot was iso- to hypo-intense in the tumor groups. Moreover, on T2-W images, a hyper-intense spot was found without obvious damage. In the animals of the PBS group, no sign of cerebral injury such as cyst formation, edema, or hemorrhage was observed (Figure 3A).

Histopathology

To evaluate the PKH+ cells tropism to the glioma mass, serial cryo-sections were evaluated using a fluorescent microscope. Clearly, the images showed the tumor area (U87-PKH+) in all the tumor-bearing groups (Figure 3B, a-c). In BMS-treated rats, BMS-PKH+ cells depicted an extensive tropism into the glioma mass and tissue cryo-sections showed no BMS-PKH+ migrating out of the glioma mass. Dual Hoechst+ and PKH+ cells were observed exclusively within the brain of BMS-treated rats. Moreover, by more careful analysis of serial cryo-sections, no BMS-PKH+ was detected in normal brain tissue or in the contralateral hemisphere of BMS-treated rats (Figure 3B, d-g).

Analysis of images related to OEC-treated animals indicated migration of OEC-PKH+ cells toward tumor mass such as BMS-PKH+. Serial cryo-sections of the OEC-treated animals revealed that OECs have substantial migration towards tumor tissue. Images suggested that many OEC cells were resettled in glioma tissue as compared to the uninvolved brain (Figure 3B, h-k). Cell tracer findings indicated that there were larger numbers (~100 cells per section) of BMS cells in the tumor area of the BMS-treated rats compared with OECs-treated rats (~76 cells per section).

In this research, PKH+ cells were infused initially via the tail vein where the results indicated the lack of these cells in the tumor tissue. Results (data not shown) showed that many PKH+ cells resettled in the lung, spleen, and liver after tail vein injection. These results highlighted the inability of tail vein transplantation of stem cells and suggested that other methods of delivery should be considered.

Intra-carotid and intracardiac methods were therefore studied. PKH+ cells were infused into the rats via the left ventricle and after the intraventricular injection the numbers of PKH+ cells were significantly increased in the tumor tissue. Moreover, our results clearly showed that intraventricular delivery of cells led to significant migration of cells towards glioma.
Figure 1: Characteristics of BMS cells. A) Cultured BMS cells observed as spindle-shaped cells. B) Flow cytometry analysis of BMS cells. C) Alizarin red staining, displayed Ca\(^{2+}\) aggregation in the extracellular matrix of induced cells (black arrowhead). D) Oil red staining exhibited lipid droplets in the cytoplasm of the induced cells (black arrowhead).

Figure 2: Characteristics of OECs. A) Cultured OECs observed as spindle-shaped cells with long processes. B-D) Immunostaining of OECs by S100β. E-G) Immunostaining of OECs by p75.
The intra-carotid route was another method that was tested. However, under intra-carotid injection, PKH+67 cells exhibited high migration and homing towards the tumor. However, during the intra-carotid procedure, the rate of animal death was very high. Due to this mortality rate, the intraventricular route was chosen as it was a safe and feasible method for cell delivery compared with the intra-carotid procedure.

H&E images related to tumor-bearing animals showed that glioma tissues were strongly stained with hematoxylin compared with the normal tissue. Neoplastic properties such as mitotic division, neoangiogenesis, and migration of cancerous cells were observed in tumor-bearing rats. Interestingly, in BMSC-treated rats, not only did BMS cells led to strongly U87 cells migration towards the normal brain parenchyma but also caused formation of metastatic masses. However, no signs of metastasis were seen in the tumor group. Also, H&E sections showed that the presence of OECs led to the reduction of tumor cell migration in the OEC-treated rats whereas, migration of U87 cells into tumor-surrounding tissues was detected in the tumor group (Figure 4 A).

For determination of neovascularization, ten H&E sections were randomly investigated, and neoangiogenesis was counted with 200 magnification. The H&E findings of the BMSC-treated animals showed that numerous neovascularization areas were formed compared with other groups. This increase of neovascularization indicated a significant difference with

Figure 3:
A: MRI of groups. a, b) The magnetic resonance images of tumor groups (white and red arrow). c) The magnetic resonance images of PBS group. B: Frozen sections of tumoral tissues were shown in the glioma-bearing animals. a) Frozen section for tracking PKH+26 tumor cells. b) Frozen section counterstained with Hoechst. c) Merged images for detection of area of tumor. d) Frozen sections of the BMS group for tracking PKH+26 cells. e) Frozen section for tracking PKH+67BMS cells. f) Frozen section counterstained with Hoechst. g) Merged images for detection of PKH+67BMS cells. h-k) Frozen section of the OEC group revealed extensive migration of PKH+67-OECs toward the tumor mass.
tumor and OEC-treated rats. While, findings related to OEC-treated rats suggested that the rate of angiogenesis was decreased compared with the tumor group, no significant difference was found between the OEC and tumor groups (Figures 4A, B).

**Immunohistochemistry**

The findings of IHC analysis depicted that Ki-67 expression was vastly different among experimental groups. Positive cells were strongly expressed in BMSC-treated rats but there was no statistical difference with the tumor group. However, the presence of OECs reduced the levels of Ki-67 in the animals. The OECs-treated rats showed fainter immunostaining than the tumor group although there was no statistical difference (Figure 5A-C and Figure 6A).

The results of endothelial-specific CD31 demonstrated that the levels of CD31 within the tumors of BMSC-treated rats were expressed in a significantly higher manner than the tumor-bearing animals. Less CD31 immunoreactivity was detected in the tumor mass of the OEC group. However, the results showed no significant difference between the OEC and tumor groups (Figure 5D-F and Figure 6A).

**Figure 4:**

**A:** H&E sections of groups.
- a) Site of tumor implantation (black arrow).
- b) Area of tumor in the BMSC group (black arrow) and metastatic masses of tumor in the BMSC group (yellow arrow).
- c) Area of tumor in the OEC group (black arrow).
- d) Angiogenesis in tumor mass in the BMSC group (black arrow).
- e) Angiogenesis in tumor mass in the OEC group (black arrow).

**B:** Angiogenesis analysis in groups.
Angiogenesis analysis between groups. Data are expressed as mean ± SEM. *: p<0.05, **: p<0.01.
Abdi Z. et al: Effects of the Two Types of Cells on Outgrowth of Glioma

IHC staining of GFAP showed poor expression in the tumor group; also, the formation of a glial edge at the tumor periphery was observed. Our findings illustrated that levels of GFAP were almost the same in BMSC-treated rats compared with the tumor group. On the contrary, not only was GFAP widely expressed in the OEC group, it also depicted a significant difference with tumor-bearing rats (Figure 5G-I and Figure 6A).

Our findings suggested that the protein level of Nestin in the OEC-treated rats was markedly upregulated as compared with other groups, which in turn represented a significant difference with other experimental groups (Figure 6A). The results of Nestin immunostaining showed no significant difference between the tumor and BMSC groups (Figure 5J-L and Figure 6A).

**Volume**

To assess the therapeutic efficacy of the PKH*<sup>67</sup>* cell, the tumor volume was determined. The mean tumor volumes in the OEC, BMSC, and tumor groups were 3.23 mm<sup>3</sup>, 5.07 mm<sup>3</sup>, and 3.26 mm<sup>3</sup>, respectively. The tumor volume analysis revealed a decrease in tumor size in rats treated with OEC. However, there was no significant difference in tumor volume between the groups treated with OECs and the tumor. In contrast, tumor volume was significantly higher in the BMSC group. Statistical analysis showed a significant growth in glioma mass in the BMSC group compared with the other experimental groups (Figure 6B).

**DISCUSSION**

This study focused on demonstrating the tropism of OECs and BMSC to glioma and generally the effects of these cells on glioma behavior in an animal model.

OECs were successfully obtained from the human olfactory bulb, suggesting that the ensheathing layers of the human olfactory bulb are a rich source of neural stem cells. The ICC results confirmed both the origin and purity of OECs in line with previous reports (17). Flow cytometry results of BMSC cells and their ability to differentiate into adipogenic and osteogenic line confirmed the origin and potential of the BMS cells (22).

In the present study, not only the migratory ability, but also the direction of the invasion of BMSC cells and OECs towards the glioma mass have been illustrated. BMS cells suggested

![Figure 5: Immunostaining of groups.](image)

A) Ki-67 positive tumor cell in the tumor area (black arrows). B) Ki-67 positive tumor cell in the tumor area in the BMSC group (black arrows). C) Ki-67 positive tumor cell in the tumor area in the OEC group (black arrows). D) CD-31 positive cells in tumor area in the tumor group (black arrows). E) Positive cells around the tumor area in the BMSC group (black arrows). F) CD-31 positive cells in the tumor area in the OEC group (black arrows). G) GFAP positive tumor cells in the tumor tissue. H) GFAP positive cells in the tumor area in the BMSC group (black arrow). I) GFAP positive cells in the tumor area in the OEC group (black arrow). J) Nestin positive tumor cell in the tumor area (black arrows). K) Nestin positive cell in the tumor area in the BMSC group (black arrows). L) Nestin positive cell in the tumor area in the OEC group (black arrows).
an active movement into the glioma microenvironment. Therefore, the findings of some researchers who reported the extensive tropism of MSCs tumor and intracranial injuries could be confirmed (16,24,30). Furthermore, excellent and directed migratory ability of human OECs towards the glioma were observed. Our findings keep in line with the data reported about the OECs ability to migrate toward pathological areas such as brain and spinal cord injuries (14). The fact that both OECs and BMS cells widely moved into the glioma tissue shows that these cells have a strong inherent and unique ability in migration and homing towards tumor.

Some of the studies indicated that the tropism of stem cells towards the tumor may be mediated by the tumor microenvironment. The tumor milieu provides some attractive chemical material such as growth factors, extracellular matrix molecules, and chemokines (6,9,30). Whereas, according to our findings, the tumor microenvironment merely could not be the cause of migration and homing of these cells. We previously illustrated that human umbilical cord mesenchymal cells showed less ability to migrate towards glioma compared with these cells (1). Thus, it seems that the extensive ability of BMS cells and OECs in localizing to glioma was a specific feature of these cells, and it could not be a function of all human cells. It is also unlikely that the migration and homing of these cells were the result of a species-specific (human stem cells-human U87 cell) tropism. Taking these findings together, the tumor microenvironment and species-specific tropism may not be the only key factors in migration and homing of the cell towards a tumor mass. The inherent nature of the cell may also play an important role in the migration mechanisms of a cell.

An important aspect of this study is the migration and homing of OECs and BMS cells towards the glioma, which supports the hypothesis that cell-based therapy could lead

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Figure 6:
A) Immunostaining analysis of groups.
Data are expressed as mean ± SEM.
*: p<0.05.
B) Volume analysis of tumor-bearing group.
Data are expressed as mean ± SEM.
*: p<0.05.
to a valuable improvement of glioma therapy. Most anti-cancer agents cannot achieve therapeutic concentrations in the brain because, there are physiologic barriers including the BBB and the high intratumor pressure (26). Therefore, only small particles such as stem cells can penetrate the BBB and reach the tumor. As a result, these cells can be efficient carriers for anti-cancer therapeutic agents’ delivery to target tissue, a critical feature of achieving therapeutic efficacy. Moreover, these cells have shown a strong chemotactic response to injuries (14,41). Thus, OECs and BMS cells may be used as tumor-selective targeting carriers for the delivery of anti-cancer agents and cytotoxic genes toward the tumor. The aforementioned findings suggest that these cells could function like “Trojan horses” for the delivery of therapeutic agents to the glioma in clinical trials.

In this study, histological slides of the brain revealed histologic features such as pleomorphism, mitotic activity, and nuclear atypia in the tumor and BMS groups. By IHC analysis and measurement of tumor volume, an association of grafted BMS cells with outgrowing glioma mass was indicated. The findings showed that BMS cells play a key role in tumor neovascularization and glioma progression. Evidence also indicates that grafted MSCs directly support the tumor vasculature and growth of tumors (28,39).

Another contribution of grafted BMS cells to tumor growth occurs through the accelerated proliferation of glioma cells. Here, it has been demonstrated that BMS cells increase the Ki-67 labeling index in BMSC-treated rats. These data are in agreement with findings from Spaeth et al.(38) which reported MSCs enhance proliferation of breast cancer cells. Furthermore, the outcome of this research illustrated that BMS cells enhance the metastatic potency of tumor. This finding suggests that grafted BMS cells may maintain the proliferative potential of tumor cells, and retain the malignant process. Recently, studies reported that MSCs promote cancer growth and metastasis (20,29). However, MSC cancer-suppressing effects have only been observed in a few reports (33,34); and most studies reported that migration of MSC leads to angiogenesis and tumor development (27,32). Moreover, extensive tropism of BMS cells to tumor was observed, which may be due to the BMS cells secreting a variety of substances such as stem cell factor, interleukins, and colony-stimulating factor- I (5,8).

In the present work, the results suggested that the presence of OECs leads to decrease of glioma tissue growth. Furthermore, the high expression of GFAP in tumor tissue was shown in OEC-treated rats, which confirmed both the migration and homing of OECs into glioma as OECs are GFAP and Nestin positive cells (31,40).

These results illustrate that the interactions between OECs and tumor cells down-regulate invasion, tumor growth, and metastasis as a mechanism of the anti-glioma effect. This can also be due to substrates and neurotrophic factors being provided by OECs that contribute to neurons regeneration and normal neuron survival. It was also observed that OECs provide least adverse reaction in the brain after transplantation, and reports have also confirmed that OECs have non-tumorigenic behavior (13,16). Thus, the multifunctional capacity of OECs has numerous positive consequences. First, OECs have a manifest tropism to the glioma mass. Second, OECs can be inherently safer than other molecular approaches that target tumor tissue, and these cells are unlikely to produce systemic toxicity. Third, OECs can be obtained from the patient. Finally, OECs can be modified in vitro to provide unique potencies and properties for GBM therapy.

With this information, future studies on migration and homing mechanisms of OECs and BMS cells towards glioma should be performed. Identification of the migratory mechanisms and the behavior of cells may lead to the application of these cells in clinical trials for the treatment of glioma.

**CONCLUSION**

In this study, OECs caused decrease of mitotic activity, as well as angiogenesis. OECs and BMS cells showed the capacity to integrate into human glioma. Moreover, these cells may be cellular carriers, which are universally applicable like “Trojan horses” for the delivery of cytotoxic genes and therapeutic agents for glioma.

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