Stem cell in alternative treatments for brain tumors: potential for gene delivery

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Abstract
Despite ongoing research efforts and attempts to bring new drugs into trial, the prognosis for brain tumors remains poor. Patients with the most common and lethal intracranial neoplasia, glioblastoma multiforme (GBM), have an average survival of one year with combination of surgical resection, radiotherapy and temozolomide. One of the main problems in the treatment of GBM is getting drugs across the blood brain barrier (BBB) efficiently. In an attempt to solve this problem, there are ongoing experimental and clinical trials to deliver drugs within stem cells. The purpose for this method is the ease by which stem cells home to the brain. This review discusses the experimental and clinical applications of stem cells for GBM. We also discuss the different properties of stem cells. This information is important to understand why one stem cell would be advantageous over another in cell therapy. We provide an overview of the different drug delivery methods, gene-based treatments and cancer vaccines for GBM, including the stem cell subset.

Keywords: Stem cells, Cell therapy, Glioblastoma, Cancer

Review
The prognosis for patients with brain tumors, in particular glioblastoma, remains poor. This poor prognosis could be explained by the hindrance to get drugs to the brain for achieving efficacious levels. Stem cells have shown tremendous promise for almost all fields of medicine including drug delivery. The application of stem cells as a cellular vehicle to deliver drugs to the brain has been noted because stem cells can cross the blood brain barrier (BBB) and can exert pathotropic effects, which attests to their ability to home to tumor sites.

Mesenchymal stem cells (MSCs) and neural stem cells (NSCs) have been used in trials to deliver prodrugs to tumors [1,2]. However, since MSCs can also support tumor growth, this represents a major disadvantage for the application of MSCs in drug delivery to target tumors [3]. Despite this disadvantage, MSCs have a unique advantage in that they can be available upon demand, generally referred to as a source of off-the-shelf cells. This ease of availability is primarily due to their ability to evade immune rejections, thereby allowing their injection across an allogeneic barrier. It is unclear if NSCs have a similar property for use on demand or if they require matching at the major histocompatibility complex class II. This review article discusses the different issues associated with the application of stem cells as vehicles of drug delivery, using glioblastoma as a representative indication.

Sources of stem cells - advantages/disadvantages
Embryonic (ESC), fetal and adult stem cells are intensely studied for their ability to differentiate into neuronal cells by in vitro methods, and are studied for their use in neurological disorders [4,5]. Each class of stem cell possesses advantages and disadvantages for treatment of neural disorders. ESCs are derived from the inner cell mass of the blastocyst and can be differentiated in vitro into all cell types. Theoretically, the intrinsic ability of ESCs to form all types of neural tissues makes them superior to other stem cells. Similarly, induced pluripotent stem cells (iPSCs), which are generated through genetic manipulation of somatic cells, have the potential to form all types of cells including those within the neuronal and glial lineages [6]. The main disadvantage of ESCs and iPSCs is their ease in spontaneous transformation. Other
issues include the ethical quandary to derive ESCs and the inefficiency to generate iPSCs.

The scientific disadvantages of ESCs and IPS led to increased interest in cell replacement strategies with adult stem cells (ASCs). More importantly, the ASCs have prospects for transplantation without ethical dilemmas. Regarding brain repair, ASCs can be effective with NSCs, MSCs, hematopoietic stem cells (HSCs) and stem cells from umbilical cord blood (UCB). Although still in experimental phase, the experimental evidence indicated that some or all of the aforementioned stem cells can differentiate into neurons and glia. There are distinct advantages of some stem cell sources over others.

NSCs are multipotent cells found within selected regions of the adult brain. NSCs can differentiate into cells of all neural lineages [7,8]. Two neurogenic areas of the brain where NSCs reside are the subventricular zone (SVZ) of the lateral ventricles and the subgranular layer of the hippocampal dentate gyrus [9]. Physiologically, NSCs are responsible for neocortical neurogenesis to help replace damaged tissue [9]. This regenerative capacitance is outweighed by the rate of neural degeneration and the amount of damaged tissues in neurodegenerative conditions. An example of this imbalance could be seen in traumatic brain injury. Subacute NSC therapy following traumatic brain injury led to cells incorporating and remaining in the tissues two weeks after transplantation [10]. The transplanted NSCs have been shown to improve the motor function of the experimental animals [10]. A major disadvantage to the utilization of NSCs is the difficulty of harvesting and isolation from an intact brain tissue. Human NSCs can be generated from differentiated ESC, iPSCs, fetal tissues sources and cadavers. None of these sources might be able to produce adequate number of NSCs for widespread clinical implementation.

MSCs are heterogeneous, multipotent cells found in several adult tissues including bone marrow (BM) and adipose. MSCs can form cells of all germ layers [11]. In BM, MSCs are found around the central sinus where they can function as "gate-keeper" cells. At this site, the MSCs contact the abluminal region of the sinus. The presence of MSCs around the central sinus is significant to the protection of BM functions [12]. The method by which MSCs protect the BM might be important to extrapolate to other organs such as neural protection effects of MSCs.

Intravenous administration of allogeneic MSCs can promote functional recovery and brain repair in experimental ischemic stroke [13]. Due to the ease of harvesting and expanding MSCs, they can be easily available from both allogeneic and autologous sources for transplantation to patients. A major advantage of MSCs to be transplanted across allogeneic barrier makes MSCs an attractive alternative for neural repair.

BM-derived HSCs were reported to have neurogenic potential [14]. HSCs are multipotent cells with their main purpose to replenish the body's immune and blood cells [15]. HSCs can be selected from the adult BM using well-defined markers. However, there are constraints for clinical application; in particular their low frequency in the BM, and their inability to be expanded. More importantly, there is no clear data that HSCs can generate neural cells. Overall, HSCs represent a less favorable source for neural repair.

UCB is a rich source of HSCs. The HSCs are functionally more immature as compared to similar cells in the adult BM [16]. MSCs can be isolated in the Wharton jelly of the cord and, to a lesser extent from UCB. If cord-derived MSCs can generate neurons or can be efficient in neural repair, this would be a major advantage because there is no risk to the donor since the cord would be otherwise discarded. Regarding hematopoietic replacement with UCB cells, there is a limitation because of the low volume of blood. This would provide inadequate number of HSCs for transplantation.

**Drug delivery - blood brain barrier (BBB)**

The brain has limited regenerative capacity sufficient only to replace modest numbers of lost cells [17,18]. The brain has shown a surprising ability for spontaneous repair in patients with stroke [19]. Accordingly, special biological protections exist to protect this vital organ, making the brain a difficult target for delivery of therapeutics. These include the hard barrier created by the skull, three meninges membranes of varying thickness/toughness, cerebrospinal fluid, and the BBB. The brain is well-protected by the BBB. The BBB is a biological fortress created primarily by a sheet of tightly knit endothelial cells. The cellular junctions are tight so that even small molecules have difficulty to cross the BBB. The BBB effectively seals off the brain from the rest of the body and enables strict selectivity in what crosses into the brain. In practice, this means that most therapeutics will not pass through the BBB. This can be problematic even for diseases affecting tissues other than the brain. For example, secondary tumors arising after breast cancer treatment can be found in the brain. This is thought to occur because most anticancer treatments do not pass through the BBB, leaving metastasized cells to reside in the brain. At a later date, they can emerge from quiescence to form deadly tumors [20].

Stem cells have been proposed as a cellular vehicle to deliver therapeutic agents to the brain. Due the difficulty of treating glioblastoma this type of cancer forms a basis to test if drugs delivered within stem cells can be effective in targeting the cancer cells (Table 1). There are a number of strategies by which the stem cells are delivered to facilitate passage of therapeutics through the
BBB (Table 2). Intranasal delivery of stem cells is not invasive and shows promise as a method of treatment. In an experimental mouse model, drug-loaded MSCs were delivered through the intranasal route and resulted in effective treatment of glioma [21,22]. Other varied approaches range from physical to chemical methods to evade the hurdles associated with the BBB to deliver drugs. An example of physical bypass of the BBB involved the use of surgical implantation of cells and drug-soaked discs designed to release the drugs in a time-dependent manner. This method has the advantage of allowing very specific drug dosing by placing the cells near or within the tumors or lesions. The disadvantage of this method is the disruption of BBB integrity caused by the surgery. In another physical approach, drug or stem cells are released outside the BBB while tight BBB junctions are loosened by ultrasound [23]. This has the advantage of temporarily affecting BBB integrity, but leaves the dosage to be empirically determined. The tight intercellular junction requires that molecules are modified for traversing the BBB. Since the junction heavily favors lipophilic molecules, drugs are chemically modified by lipidation and glycosylation [24]. An alternative approach is to load the drugs in liposomes for passage through the BBB [25,26]. These strategies would allow for control of the drugs passing into the brain. However, increased amount of drugs into the brain can be hazardous because passage out of the BBB would be subjected to the same constraints as passage into the BBB. Thus, there will be an accumulation of drug metabolites that could lead to toxicity over time.

In the future, investigation into novel pathways regulating BBB permeability may lead to new strategies for modulating the permeability. There are some indications that BBB permeability is subject to alterations in some diseases. For example, neurodegenerative diseases such as neuropsychiatric systemic lupus erythematosus and Parkinson’s disease have been reported to show increased permeability of the BBB [37,38]. The BBB can also be altered under non-disease conditions to facilitate the immune response. This could occur by the involvement of components associated with the BBB such as pericytes. The pericytes can respond to changes in the microenvironment by changing their structures and shapes to modulate BBB permeability [39]. If it were possible to mimic the pathways responsible for opening the BBB for immune response, methods could be developed to trick the BBB into temporarily opening up to allow delivery of cellular therapies.

### Viral gene therapy

This section discusses approaches involved in the use of engineered viruses to deliver enzymes that can convert prodrugs into toxic metabolites. Viruses preferentially infect rapidly dividing tumor cells. Figure 1 shows how virus-containing genes could be packaged for cancer therapy. Here, the non-dividing nature of neural cells would be spared by the infection to deliver the virus. A commonly used method is to combine the gene delivery system with a gene that expresses the enzyme, thymidine kinase, followed by ganciclovir treatment. The infected cells are able to translate the enzyme thymidine kinase, which phosphorylates ganciclovir to ganciclovir triphosphate. As a result, only the infected, rapidly dividing cells, are killed by the toxic metabolites and apoptosis is induced both in transduced cells and also adjacent dividing cells (“bystander effect”) [40]. Numerous preclinical and phase I/II studies have investigated this procedure and showed promising results. In an in vivo study using a rat model of cerebral glioma, fibroblasts carrying the herpes simplex virus type 1 thymidine kinase (HSV-tk) gene through retroviral vector were stereotactically injected and subsequently treated with ganciclovir (GCV) to produce a therapeutic response [41]. Clinically, a pilot trial using HSV-tk-containing adenoviruses (AdV-tk) and GCV in 13 patients with recurrent malignant brain tumors showed acceptable toxicity [42].

A phase Ib trial with relatively few patients (n = 12) who were newly diagnosed with malignant glioma received AdV-tk via tumor bed injection at time of surgery. This was followed by valaciclovir with overlapping radiation therapy, then treatment with the chemotherapeutic temozolomide. The treatment showed no significant toxicity and resulted in survival of 33% at 2 years and 25% at 3 years. The patient-reported quality of life was stable or improved after treatment. Significant T-cell inflammatory infiltrate was found in four re-resected tumors, implying long-lasting immune stimulation [43].
A phase II trial, which utilized an adenoviral vector to treat initial and recurrent high-grade gliomas showed a significant improvement in survival when compared with historical controls. However, the subsequent randomized phase II trials did not show any statistically significant improvement in survival [44]. Based on these conflicting results in preclinical and phase 1/II studies, a phase III study was conducted in adults with previously untreated GBM [45]. This trial utilized HSV-tk and GCV gene therapy as an adjuvant to surgical resection and radiation. The trial involved 248 patients that received either surgical resection and radiotherapy, or surgical resection and radiotherapy plus adjuvant gene therapy at the time of surgery. The experimental treatment was confirmed to be safe; however no difference was noted for overall survival or disease progression.

A more recent phase III study (n = 250) on the clinical use of Adenovirus-mediated gene therapy with sitimagene ceradenovec (replication-deficient adenovirus) followed by intravenous GCV (ASPECT trial) has been conducted [46]. The study compared standard of care (surgical resection followed by radiotherapy and chemotherapy) with perilesional injection of sitimagene ceradenovec followed by GCV. The median time to death or re-intervention was longer in the experimental group (p = 0.006) and, in a subgroup of patients with non-methylated methylguanine-DNA methyltransferase (MGMT) the hazard ratio (HR) was 1.72 (95% CI 1.15-2.56; p = 0.008). However, there was no difference between the two groups in terms of overall survival. One possible explanation for the discordant results observed in the studies is the variable gene delivery and transduction rate in the tumor cells, which has been rarely measured. This observation encourages further efforts focused on the optimization of drug delivery administration [47].

Use of stem cells in glioblastoma treatment

Neoplasias, including gliomas, are heterogeneous diseases. There is overwhelming in vitro and in vivo evidence that a subpopulation of cancer stem cells (CSCs) initiates and sustains tumor growth, resulting in tumor masses with heterogeneous malignant cells [48]. It has also been suggested that a subpopulation of glioma CSCs (gCSCs) may be responsible for the resistance to standard therapy. Eradication of the gCSCs could lead to significant improvement in patients’ outcomes [49].

Targeting cancer stem cells (CSCs)

Targeting CSCs could be most efficient to prolong the survival of cancer patients. Several microRNAs, such as miRNA-145 and oncomiR-138, have been identified for this purpose. The oncomiR-138 is considered to be a molecular signature of gCSC and has been targeted by functional inhibition in vitro, resulting in decreased tumorigenesis and impairment of gCSCs growth [50]. In vitro studies showed a tumor suppressive effect of miRNA-145 in glioblastoma [51]. The suppressive effects of miR-145 occur by its ability to decrease the expression of stem cell-linked genes within the CD133 expressing CSC-like cells [51]. Ectopic delivery of miR-145, in combination with radiotherapy and chemotherapy, improved survival in an experimental model of glioblastoma [51]. The inhibition of the Notch pathway through gamma-secretase inhibitors has also been experimentally studied as a targeted treatment for gliomas [52].
This was performed by the implantation of a drug-impregnated, polymer bead delivery system. This treatment method blocked tumor growth and prolonged the survival of a small cohort of mice.

**Stem cells in gene and drug delivery**

Studies are proposed with NSCs as a possible cellular vehicle to deliver suicide genes to tumors. The advantage of this approach is that NSCs are tumor tropic. The tropic effects are likely facilitated by the production of several chemoattractants by glioma cells, such SCF-1 or MCP-1 [53]. Tables 1 and 2 show a snapshot of how drugs can be delivered to tumors with the use of stem cells. Also shown in Table 2 are the different approaches to find the most efficient method to deliver the drug-loaded stem cells to the brain. There are few reports describing the use of NSCs to deliver the enzyme cytosine deaminase (CD), followed by treatment with 5-Flucytosine (5-FC). 5-FC can be converted by CD to the cytotoxic compound 5-fluorouracil (5-FU), which selectively targets tumor cells, with minimal toxicity to the surrounding healthy tissues [1]. Clinical trials using a similar approach are ongoing [54]. Programmed self-destructive NSCs have also been used as a delivery tool for pH-sensitive Mesoporous Silica Nanoparticles-doxorubicin (MSN-dox); the NSCs migrate to the tumor site, eventually undergo apoptosis and release the MSN-dx to the surrounding gCSCs. [55]. MSC have been used in several experimental models as an efficacious tool for drug delivery. Delivery of soluble (s)-TRAIL via MSCs has been shown to induce apoptosis in glioma. Mice were implanted with a mix of gCSCs and S-TRAIL-expressing MSCs or a mix of gCSCs with GFP-expressing MSCs as control. Real time imaging showed a significant reduction in tumor burden in animals implanted with MSCs expressing s-TRAIL as compared to controls [56].

Recent studies showed a role for miR-9 in the expression of the drug efflux transporter, P-glycoprotein in TMZ-resistant GBM cells [2]. MSCs loaded with anti-miR-9 were able to deliver the drug through exosomes thereby re-sensitizing the GBM cells to TMZ [2]. MSCs were used to deliver the CD suicide gene in vivo [27]. Intracerebral inoculation of engineered MSCs after glioblastoma surgical resection resulted in a curative outcome in a significant number of mice [27].

Marrow-isolated adult multilineage inducible (MIAMI) cells, which are believed to be a subset of MSCs, have been used to deliver lipid nanocapsule loaded with an organometallic complex [28]. In vitro and in vivo studies indicated that this type of drug delivery resulted in cytotoxic effect on the glioma cells [28]. MSCs, engineered to express a single-chain antibody to EGFRvIII, were co-injected with EGFRvIII (+) glioblastoma in a xenograft model. The MSCs-EGFRvIII showed significant increase in the survival of the mice as compared to the controls injected with glioma cells alone [57].

**Oncolytic viruses**

Another possible approach to treat brain tumors involves the use of oncolytic viruses. These viruses preserve the ability to replicate, selectively amplifying in cancer cells to cause cytotoxicity. The oncolytic virus can also activate and stimulate the natural immune response to target infected cells. One example of a widely used oncolytic virus is herpes simplex virus (HSV), primarily because of its cytotoxicity and ability to induce a strong immune response [58].

Several phase I/II studies investigated this technique using various engineered viruses, such as HSV, adenovirus, retrovirus and others, and showed overall safety, low rate of complications, and promising effects [59-61]. Cheema et al. combined oHSV and IL-12 to enhance the immune response and at the same time prevent angiogenesis [62]. The transgenic oHSV (G47Δ-miL12) showed a significant increase in survival, decreased neovascularization, decreased VEGF expression, and increased production of angiostatic IP-10 (CXCL10). The efficacy of the transgenic oHSV was markedly reduced in athymic mice, suggesting a direct effect of T-cell stimulation with IL-12.

**Anti-cancer vaccines and stem cells**

The use of anti-cancer vaccines has been studied for the treatment of several solid tumors, resulting in discordant results [63]. These therapies re-sensitize and enhance the natural immune response of the host against tumor cells. In the absence of the vaccine the tumors can evade the immune response.

One method involves the use of dendritic cells (DCs) from patients for sensitization to the tumor’s unique antigens. The DCs are then reinfused in the patient, where they can activate cytotoxic T-cells against the tumor cells. Several phase I and II studies showed feasibility and safety of this approach in the treatment of gliomas, with some encouraging results [64-66]. Another easier approach to cancer vaccine is the use of formalin fixed tumor cells, collected at tumor’s resection time, as a stimulant for immune response. This approach has been studied in phase I/II studies with promising results [67]. The combination of cancer vaccine with EGFR inhibitors has also been explored in a phase II study, showing safety and increased survival [68].

Targeting the subpopulation of glioma Cancer Stem Cells (gCSCs) through immunotherapy could lead to the eradication of gCSCs. This particular subpopulation of cells is therefore the ideal candidate for vaccine therapy. Ji et al. took advantage of CD133, a proposed marker of gCSCs, as a target to stimulate cytotoxic T-cells (CTLs)
Peptide-specific CD8+ CTLs from normal donors were generated and pulsed with autologous DCs. The CTLs efficiently recognized the CD133 epitopes and were specifically able to lyse CD133+ gCSCs in vivo.

Chimeric antigen receptor
Chimeric antigen receptor (CAR) T-cells were first developed in an attempt to expand the therapeutic potential of effector lymphocytes in adoptive T-cell transfer, a term first coined in the 1950s [70]. In a landmark study, the single-chain of an Fv (scFv) antibody molecule was fused to the γ chain of the Fc receptor or to the ζ of the CD3 complex, creating T cells with antibody type specificity and subsequent IL-2 signaling leading to target cell lysis [71]. This target-binding site displays an affinity much higher than TCRs, and in addition is MHC independent, avoiding tumor escape mechanisms secondary to MHC loss variants. The advancements in the field to characterize “designer lymphocytes” provide a scaffold for cell-based immunotherapies. Here we discuss the CAR T-cells as a method of immunotherapy for brain tumors because these cells show enhanced efficiency to enter the brain (Figure 2). Indeed, there are ongoing studies to use CAR T-cells to EGFR, a common receptor on glioblastoma [72].

Over the years, CARs have been engineered and manipulated to achieve more targeted and potent effects. The need for this became clear upon further understanding of activating ligands on antigen presenting cells such as CD80 and CD86 that bind to the co-stimulatory receptors found on T-cells including, but not limited to CD28 [73]. Further focus turned towards incorporating co-stimulatory signals into the domain in order to prevent T-cell apoptosis or anergy [74,75]. This has otherwise led to the development of second and third generation CARs with greater effects than previous generations.

The advantages of CAR therapy are many including HLA-independent recognition of target antigens and the ability to rapidly deliver a large population of tumor antigen-specific T-cells. However, there are important disadvantages that must be addressed, including ‘on target/off tumor’ effects, explained by antigen similarities that may be shared by normal cells and tumor cells, and cytokine-release syndrome.

Cytokine-release syndrome is driven by pro-inflammatory cytokines such as IFN-γ, TNF-α and IL-2 [76,77] or more recently described, IL-6 [78]. The effects of IL-6 was demonstrated with studies using tocilizumab (anti IL-6 receptor monoclonal antibody) in glucocorticoid resistant GvHD [79]. Cytokine-release syndrome seems to be related, however with T-cell expansion as patients present clinically with fever, variable degrees of myalgias, nausea and anorexia and with complications that ranged from hemodynamic or respiratory instability [76].

Medically, there are clinical manifestations that limit the use of CAR therapy. CAR therapy is contingent on the immunogenicity of the target cells. The growing field of cancer stem cells (CSCs) showed that they should be the target cells. The identification of markers on CSCs

![Figure 2: Relative efficiency of drug delivery in cells or alone.](image)
is a subject of investigation, lead to questions. How immunogenic are CSCs? There are also concerns to the fate of CAR cells upon entering the tumor microenvironment. Lastly, most data pertains to hematological malignancies; yet some data, although limited with modest or null results, have been done on solid tumors, including neuroblastomas [80].

There are recent advances with promising results in CAR therapy. The mechanisms behind cytokine-release syndrome and its associated neurotoxicities must be addressed. These limitations may further be explored with future studies that include greater power. This may pave the way for engineering of other cells, as CAR modifiable cells are not limited to T-cells, but also include, but are not limited to NK cells, iNK T cells.

**Conclusion**

Glioblastoma’s aggressive behavior seems to be determined by the extreme heterogeneity of this tumor, which can be sustained by gCSCs. The CSCs can generate, re-generate and maintain tumor growth. The challenges related to the treatment of glioblastoma are represented in the first place by the BBB, which physically isolates the brain from the rest of the body, causing difficult delivery of chemotherapeutics and reducing the access of cells of the immune system. Figure 2 shows the relative efficiency of delivering drugs directly or through stem cells. The stem cells show promise to enter the brain with higher efficiency to directly deliver the drugs to brain.

Glioblastoma is indeed often able to evade the natural host’s immune response. Studies focusing on successful delivery of therapeutics have been conducted to determine if the permeability of BBB can be modulated to enhance the transport of drugs to the brain. Further studies focused on the regulatory pathways of the BBB’s permeability could allow delivery of targeted therapies in specific time frames.

Different types of stem cells have been studied as a method of optimal delivery of suicide genes and drugs, due to their unique ability to migrate to the tumor bed with adequate specificity (Table 3). MSCs can be relatively easily harvested and expanded, however concerns regarding the potential for transformation needs to be thoroughly addressed [81].

Gene therapy has been broadly studied for glioblastoma, and various techniques for gene delivery involving the use of stem cells as transporters, have been investigated with promising results in vitro and in preclinical studies; unfortunately phase III clinical trials failed to demonstrate a clear advantage of experimental therapies in terms of OS. This could be explained by low delivery rate and/or inconsistent levels of transductions of the drugs, therefore further research focused on improvement of delivery methods could potentially bring significant improvements to glioblastoma gene-based treatments [47].

Finally, great hope is represented by strategies based on the enhancement of natural host immune response, which is frequently evaded by the tumor: anti-cancer vaccines targeting gCSCs would theoretically allow the eradication of this cell subpopulation, likely responsible of recurrence and resistance to chemotherapy. This study addresses the use of CAR T-cells, which shows promise. Its inclusion in this brief review is mainly due to the relative ease to enter the brain [72].

The use of MSCs or NSCs as a delivery tool for either specific genes or drugs has been explored in vitro and in vivo showing promising results in terms of tumor cells cytotoxicity and increased survival in animal models. These methods clearly offer enormous advantages for the potential treatment of brain tumors, being able of targeting almost uniquely cancer cells. Few clinical trials addressing the feasibility and safety of these approaches are ongoing (Table 2).

In conclusion, as an extreme heterogeneous disease, complicated by the peculiar localization in a selectively protected environment, glioblastoma is a disease that needs to be approached from different angles, keeping in mind the unique limits and potentials of the organ from which it raises from. These features, even though extremely challenging, offer potential starting points for future research.

**Competing interest**

The authors declare that they have no competing interests.

**Authors’ contributions**

WM prepared the section on gene therapy, summary, conclusion and assembled the different parts of the review; SLJG prepared the section on the advantages and disadvantages of stem cell types; RM prepared the section on drug delivery; GRN wrote sections of the review on immunology; PR

### Table 3 Cancer targeting agents delivered by stem cells for glioblastoma

| Drug/gene Stem cell | Targeting method | References |
|---------------------|------------------|------------|
| Cytosine deaminase (CD) | NSCs | Indirect via conversion of prodrug, 5-fluorocytosine (5-FC) | [36] |
| CD | MSCs | Indirect, via conversion of the prodrug, 5-FC | [27] |
| MSN-dox | NSCs | Direct | [55] |
| Soluble TRAIL | MSCs | Direct | [56] |
| miR-9 | MSCs | Direct | [2] |
| Fc-diOH-LNC | MIAMls | Direct | [28] |

Shown are representatives methods by which drug/gene/RNA can be used in stem cell delivery system for the treatment of glioblastoma. Direct method indicates that the stem cells release the drug, which interacts with the cancer cells for cytotoxic effects. Indirect effects are indicated when an enzyme is delivered in the stem cells for the local conversion of a prodrug to its active form.
managed the review, corrected the text and made the final edits for submission. All authors read and approved the final manuscript.

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