Therapeutic nanomedicine for brain cancer

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Abstract

Malignant brain cancer treatment is limited by a number of barriers, including the blood–brain barrier, transport within the brain interstitium, difficulties in delivering therapeutics specifically to tumor cells, the highly invasive quality of gliomas and drug resistance. As a result, the prognosis for patients with high-grade gliomas is poor and has improved little in recent years. Nanomedicine approaches have been developed in the laboratory, with some technologies being translated to the clinic, in order to address these needs. This review discusses the obstacles to effective treatment that are currently faced in the field, as well as various nanomedicine techniques that have been used or are being explored to overcome them, with a focus on liposomal and polymeric nanoparticles.

Current state of malignant brain cancer

Nearly 25,000 new cases of malignant brain cancer are estimated to be diagnosed in the USA each year, accounting for close to 15,000 deaths [1]. This review focuses on gliomas, the three major classes of which according to WHO classification are astrocytomas, oligodendrogliomas and mixed gliomas (oligoastrocytomas) [2]. The most common of these are astrocytomas, which the WHO separates into various grades of severity based on morphological abnormalities, increased angiogenesis and proliferation, and the presence of necroses. Malignant or high-grade astrocytomas include the grade III anaplastic astrocytomas and grade IV glioblastoma (GB) and are the most common gliomas. Generally, malignant astrocytomas are treated by surgical resection, chemotherapy and radiotherapy, with the goal of prolonging survival time and improving quality of life rather than curing the disease. The median survival for patients with grade III and IV glioma is 2–3 years and 14–15 months [3,4], respectively, prompting the need for novel treatments that can promote a better outcome.

The field of nanomedicine has opened opportunities for improved brain cancer diagnosis and treatment. Nanoparticulate systems are already in use in the clinic for enhancing treatment of various other cancers. Because of the wide variety of biocompatible materials being researched, the range of nanocarrier compositions and design strategies is very wide. While many of these methods are still experimental, nanoparticles have shown great promise for improving delivery of imaging agents and chemotherapeutics. Nanoparticles have been a
major focus of research in the drug delivery field because of a wide range of advantages afforded by their small size and other properties. For the purposes of this review, the term ‘nanoparticle’ will be used in reference to any particle of any shape and material below 1 μm in diameter, general types and properties of which will be described briefly. Barriers to effective brain cancer treatment (Table 1; Figure 1) will be discussed, as well as strategies currently under investigation to overcome them.

**Types of nanoparticles & general properties**

Nanoparticles can be fabricated using a wide range of different materials. Synthetic polymers, such as poly(glycolic acid), poly(lactic acid) (PLA), and the copolymer poly(lactic-co-glycolic acid) (PLGA), are biocompatible in a number of systems and are hydrolytically degradable, leading to their use in encapsulating and delivering many types of therapeutic molecules, including conventional small-molecule drugs and large biological macromolecules such as proteins and nucleic acids [5]. Other degradable polymers, including chitosan [6, 7], poly(beta-amino esters) [8, 9], poly(amidoamines) [10], and many other cationic polymers have also been used to deliver therapeutics to brain cancer models. These positively charged polymers are studied particularly for delivery of nucleic acids, because their cationic character allows them to form nanocomplexes with negatively charged DNA or RNA. While degradable cationic polymers are being studied as an alternative to virus-mediated gene transfer, viruses are also under investigation for their potential to treat brain cancer. Amphiphilic lipids can also self-assemble to form liposomes, artificial vesicles that can be used to encapsulate and deliver either hydrophobic or hydrophilic molecules and whose size can be controlled by the size and properties of the constituent lipids [11]. For example, the chemotherapeutic doxorubicin, with a circulation time of 10–15 min as the free drug, can be encapsulated in liposomes to increase this time to over 2 h [12].

Inorganic nanoparticles can also be used for imaging as well as drug delivery purposes, with advantages of highly tunable and reproducible synthesis processes. Injectable superparamagnetic iron oxide (SPIO) nanoparticles can provide contrast for MRI [13] while other metallic nano-particles, such as materials like gold, can be used to carry a conjugated drug or be used for photo-thermal therapy [14, 15]. Mesoporous silicon particles can be loaded with therapeutics or other smaller, drug-loaded nanoparticles for multi-stage delivery or imaging systems [16, 17].

The surface of any of these types of nano-particles can be modified to change their transport properties and biological responses to them. One common modification is PEGylation, coating polyethylene glycol onto a surface via physical interaction or covalent conjugation. PEGylation creates a hydrophilic and non-charged surface, which can prevent early clearance of nanoparticles and thereby increase their circulation time [18]. For example, liposomes are often cleared quickly from circulation by the mononuclear phagocyte system [19, 20], but their half-life in circulation increases from 2 to 40–71 h when the liposomes are PEGylated [12]. In addition, nanoparticle size, an important property in determining drug delivery efficacy, can often be varied during particle fabrication. A particle’s size affects its non-specific uptake by cells and clearance as well as its ability to deliver intracellular drugs to target cells. Intracellular delivery remains a general challenge in drug delivery because of the plasma membrane that prevents penetration of many drugs and downstream barriers that cause degradation of the drug or prevent it from arriving at the necessary location for activity [21]. The use of a wide variety of drug delivery systems has been explored to increase intracellular drug concentration or to avoid drug efflux as a mechanism of resistance [22]. In particular, the small size of nanoparticles has been demonstrated to allow cellular uptake [1, 23, 24]. Although uptake is also affected by other factors, including
particle composition and surface properties [25,26], particle sizes between 5–500 nm have demonstrated uptake in different cell types and delivery systems, further emphasizing the utility of nanomedicine in cancer treatment.

Another general barrier to effective drug delivery to tumors is the problem of achieving high drug concentration at the site of interest. Concentration of a drug at a tumor site is often limited by short circulation time of the drug due to nonspecific uptake or accumulation in healthy tissue, degradation, and excretion. Nanoparticulate systems have been used to improve the half-life of common tumor drugs like doxorubicin.

**Barriers to delivery to the brain**

**The blood–brain barrier**

While radiotherapy has been shown to improve prognosis in malignant glioma, the benefits of chemotherapy are controversial, with clinical trials showing wide variation in whether or not, and to what degree, chemotherapy confers a survival benefit in these patients [27,28]. A challenge specific to delivery of high concentrations of drug to the brain is the presence of the blood–brain barrier (BBB) [29], the arrangement of endothelial cells in capillaries of the brain that ensures particularly tight regulation of transport between the bloodstream and brain tissue. Several methods have been explored to allow transport of a therapeutic across the BBB, each with different advantages and safety concerns [30], some of which will be discussed broadly below. However, as in cancers of other systems, GB tumors are fed by a ‘leaky’ vasculature, an abnormal and disorganized network of blood vessels formed by the high rate of angiogenesis that characterizes high-grade malignant tumors [31]. The discovery that molecules and small particles can more easily extravasate from tumor vasculature and are returned inefficiently to circulation due to poor lymphatic drainage was a major finding in cancer therapy research and was coined the enhanced permeation and retention (EPR) effect [32].

In addition to carrying intracellular drugs and increasing cargo half-life, nanoparticles have been of increasing interest to the cancer drug delivery field for their ability to take advantage of the EPR effect. Particles less than approximately 100 nm in diameter extravasate preferentially from tumor vasculature, due to increased vascular permeability compared with normal tissue, and particles greater than approximately 20 nm are retained in the tumor tissue rather than returning freely to the circulation [33]. Larger particles are unable to leak out of the disorganized blood vessels that supply tumors, while smaller ones are less effectively retained. This effect depends primarily on the physical properties of the nanoparticles – particularly size and the ability to remain in circulation long enough to extravasate into a tumor – but it is limited by a lack of specificity for particular tissues.

The question of whether or not – and to what degree – the BBB is a major obstacle for drug delivery to brain tumors has not seen consensus in the literature over the past decades, although various studies using conventional chemo-therapeutics have found high variability or often only minimal enhanced drug retention at the glioma site over other tissues [34]. Some researchers have found that the EPR effect can allow for accumulation in the tumor via the porous brain tumor–blood barrier (BTBB) and have reported accumulation of imaging agents or particles on the order of 100 nm in diameter [35]. Many others have found that this passive targeting alone cannot overcome the BBB or does so to a degree that is insufficient for effective treatment, showing that only smaller particles (approximately 20 nm) or smaller (<12 nm) [36] are able to cross the BTBB [37,38]. The tumor vasculature is inhomogeneous [39], and some studies show that the BBB remains largely intact at tumor margins despite being more permeable within the bulk of the tumor, which hinders delivery of drugs to the tumor cells most directly responsible for invasion and migration; in addition,
increased interstitial pressure within tumors tends to oppose passive diffusion out of the circulation and into the tumor tissue [40]. As a result, although studies have found that small nanoparticles and drugs are able to take advantage of the leaky tumor vasculature [41], various methods to further improve penetration of the BTBB must also explored.

One method to improve delivery is the use of BBB disruption to further increase the permeability of these vessels by increasing the local osmotic pressure within brain vasculature using hyperosmotic agents [29]. However, this method is known to have significant patient-to-patient variability, which impedes widespread and consistent use [42]. Furthermore, all of these BBB disruption methods cause decreased integrity of the entire BBB, not specifically the vasculature of the tumor, which could adversely affect healthy brain tissue. Aside from non-specific delivery of the chemotherapeutic to healthy tissue, which can lead to unwanted toxicity and dose-limiting side effects, BBB disruption can also allow leakage of unwanted molecules from the circulation into the brain through the globally compromised BBB [37]. To avoid disrupting the entire BBB, various strategies have been employed to penetrate the BBB via the nanoparticle drug carriers themselves.

**BBB-penetrating nanoparticles**

The LDL receptors on endothelial cells of the BBB can facilitate uptake of nanoparticles coated with ligands for the LDL receptor. Xin *et al.* showed that Angiopep-coated PEG–poly(caprolactone) nanoparticles could accumulate in the tumor bed in vivo due to the passive EPR effect as well as to the active targeting through Angiopep [43]. In addition, apolipoprotein-coated particles both in vitro [44], using apoA-I-coated protamine/oligonucleotide nanoparticles, and in vivo [45], using apoE-coated serum albumin nanoparticles are taken up by binding to the LDL receptor. By coating the particles with an agent that causes absorptive uptake, this study demonstrated uptake of drug-containing nanoparticles by the BBB endothelium via adsorption by the cells, rather than increased brain penetration via increasing gaps between endothelial cells in the entire brain. Using a similar principle, Kreuter *et al.* developed poly(butyl cyanoacrylate) nanoparticles as carriers for a peptide that normally could not cross the BBB in any measurable amount. By coating the particles with a surfactant that caused apolipoprotein deposition onto the nanoparticles once in the plasma, they achieved uptake of their particles by absorptive uptake [46]. The use of polysorbate 80 [47] and other surfactant coatings, such as poloxamer 188 [48,49] and Tween® 80 [50] has also been shown to cause high uptake by BBB endothelial cells in models in vitro and led to higher accumulation in the brain in vivo, with varying degrees of toxicity depending on the particular system studied. Liposomes have also been demonstrated to increase BBB penetration [51]. Other lipid-based nanoparticles have been reported to have BBB-penetrating properties as well as the ability to take advantage of the EPR effect in tumor models [52].

By including the BBB-penetrating agent as part of the delivery vehicle itself, inducing endocytosis by endothelial cells rather than physical disruption of the endothelial layer, the use of nanocarriers can reduce the abnormal passage of other molecules into and out of the vasculature of the brain. Although with BBB-penetrating nanoparticles, nanomedicines may no longer be restricted in size by the gaps between endothelial cells, particle size is also an important factor in cellular uptake. Endocytosis, whether clathrin- or caveolae-dependent or -independent, generally involves the formation of vesicles less than 150 nm [53]. In order to either exploit the EPR effect or induce internalization by endothelial cells, particles being studied as potential drug carriers to brain tumors are generally less than 150 nm in diameter.
Surface-modified nanoparticles for BBB penetration

Another advantage of nanoparticle drug carriers in BBB penetration is their chemical versatility. Aside from surface coatings to increase uptake, such as surfactants or stealth coatings such as PEG, ligands and other biological or chemical moieties can be conjugated to the surface to promote active uptake by cells on the luminal side, trafficking of the particle through the endothelial cell, and then exocytosis into the brain tissue. As an example of the various functions that can be designed into nanoparticle systems, PLA nanoparticles were surface-modified with PEG for stability and with cationic serum albumin for increased circulation time. The cationized albumin was able to facilitate interaction of the particles with brain endothelial cells to promote uptake with little to no toxicity observed [54]. Because viruses are also essentially nano-carriers that have evolved an efficient ability to cross cellular barriers, including the BBB, virally derived ligands have also been used to penetrate the BBB, such as the HIV Tat peptide. Polymeric core/shell nanoparticles, made of amphiphilic polymers that form micellar structures, were conjugated to Tat peptide, allowing them to cross the BBB into the brain [55]. Qin et al. used Tat-conjugated cholesterol to formulate liposomes that showed the ability to transcytose through brain capillary endothelial cells and accumulate in the brain [56]. In another study, poly(amidoamine) dendrimers were conjugated to a peptide derived from the rabies virus glycoprotein, RVG29, through a PEG linker and conjugated with DNA to form nanoparticles. The RVG29-modified DNA-containing nano-particles accumulated in the brain significantly more efficiently than unmodified nanoparticles (Figure 2) [57]. Other researchers have taken advantage of toxins that increase vascular permeability, such as the diphtheria toxin. By conjugating a mutated ligand for the diphtheria toxin receptor (CRM197) to liposomes, Van Rooy et al. were able to show in vitro that CRM197-modified particles localized to the endothelium but that other ligands, such as RI7217, an antibody binding to the transferrin receptor (TfR), had the greatest effect in vivo [58]. Specificity can also be built into the delivery system by use of specific ligands that promote receptor-mediated endocytosis. Insulin is transported across the BBB from the circulation by receptor-mediated transcytosis [59], and an antibody to the insulin receptor was taken up effectively into the brain in vivo in a primate model [60]. Ulbrich et al. conjugated drug-loaded human serum albumin nanoparticles to antibodies against insulin receptor and were able to achieve uptake into the brain in a mouse model [61]. Another receptor commonly studied for active transport across the BBB is the TfR, which is necessary for transport of iron into the brain and was used to deliver similar serum albumin nanoparticles into the brain [62]. In particular, its expression is high in rapidly dividing cell populations [63], which, while not seen only in malignancies, does have some degree of specificity for rapidly growing tumors. Although TfR is expressed in other tissues such as the liver and bone marrow [64], it is still under active investigation for transport across the BBB. Polyester nanoparticles loaded with anti-cancer drugs like taxols, based on materials such as PEG-conjugated PLA or PLGA [65,66], were able to achieve higher accumulation in brain endothelial cells when conjugated to transferrin. Like transferrin, the folate receptor for transport of folic acid is also upregulated in rapidly dividing cells, including those in malignant brain cancer [67].

Nanoparticle movement within brain tissue

Once a drug passes through the BBTB and into the brain tissue, other barriers still remain. Because surgical resection is normally the initial step in treatment, local delivery of a chemo therapeutic to the site of the tumor is possible and may be a reasonable way to bypass the delivery bottleneck of the BBB. However, high-grade gliomas are highly invasive, and cancer cells are often found extending out of the main tumor bulk [68]. Not only does this complicate surgical resection, as it is difficult or impossible for the surgeon to visualize all
tumor cells, let alone remove them without extensive damage to the surrounding brain tissue, but some controversy still exists over the extent of benefit from resection in some cases of malignant glioma [69–71]. As a result, the ability of any drug or drug carrier to diffuse through the brain tissue and affect all relevant cells is imperative for effective treatment. The majority of GB recurrences are seen at or adjacent to the original tumor site [72]. While this indicates that local delivery may be sufficient in many cases as long as the drug or drugs can affect all tumor cells in the area, it also shows that current therapies are thus far unable to kill all tumorigenic cells at the tumor site. A study by Kroin et al., which used local delivery of cisplatin via small cannulas, found that multiple cannulas would be needed to maintain a therapeutic drug concentration for tumors greater than 1 cm; for tumors greater than 2 cm, local delivery would not be sufficient [73]. While this is not the only local delivery system that has been studied, the study’s conclusion raises the concern that most drugs under investigation do not diffuse freely enough through the brain tissue for full efficacy. Furthermore, low diffusion following local delivery can result in high local toxicity at the delivery site. As a result, studies over the last decade have generally used one of the following methods to enhance interstitial movement of therapeutics through the brain.

### Nanoparticle diffusion within tissue

Several strategies have been employed to overcome the poor movement of therapeutics within the brain interstitium. Initially, attempts to better understand the dimensions of the extra-cellular space, modeled as pores through which particles can diffuse in the brain, illustrated that particles must be approximately 30–70 nm in diameter or smaller to move through the brain interstitium [74]. In one study, PLA–PEG nanoparticles were loaded with aclarubicin and coated with cationic albumin [75]. By keeping particles within the approximate range stated above, the authors demonstrated improved survival in a rat glioma model using this nano-particle formulation, suggesting that their system was able to effectively deliver particles through the disrupted BBB and move through the tumor interstitium.

However, it has since become clear that, while size is an important factor, nanoparticle surface properties can also alter diffusion through the brain. Nance et al. found that polystyrene nanoparticles (>100 nm) and quantum dots (35 nm) diffused very slowly through brain interstitium when unmodified or coated sparsely with PEG; however, a dense layer of PEG allowed significantly increased movement through the brain with particles of at least 114 nm in hydrodynamic diameter [76]. Therefore, while smaller particles will tend to allow greater diffusion than larger ones, the upper limit of nanoparticle size can be extended with surface modifications that increase the particles’ diffusivity, thereby increasing the capacity of the particles as drug-loaded nanodevices.

### Convective transport of drugs & nanoparticles

Another way to increase transport through tissues is to rely on fluid convection during local delivery, or bulk flow, as well as diffusion. A hydrostatic pressure gradient created in the tissue can allow large molecules or particles to be carried much more quickly throughout the tissue. For instance, the well-established local delivery system of the 1,3-bis(2-chloroethyl)-1-nitrosourea (BCNU)-loaded Gliadel® wafer (Arbor Pharmaceuticals, LLC, GA, USA) is known to be limited in part because the low diffusion of BCNU through the tumor tissue cannot overcome the rate of drug clearance; however, this system does demonstrate higher initial penetration than in later days, an unexpected finding that was thought to result from convection due to transient local edema following surgery and placement of the wafer [77]. Bobo et al. demonstrated that continual infusion of a drug solution into feline brains via cannulae over the course of approximately 2–3 h, establishing high fluid flow, caused greater penetration of their drug into the brain tissue as well as more
homogeneous drug distribution [78]. At the infusion rates used in this study, as well as in other studies using convection-enhanced delivery (CED), the rate of diffusion is very slow or negligible compared with that of convection by bulk flow.

In a study by Sawyer et al., PLGA nano-particles between approximately 90–120 nm mean diameter were loaded with camptothecin and injected intracranially into tumor-bearing rats. The median survival time in rats treated with camptothecin nanoparticles delivered by CED was longer than in rats treated with the same dose of unencapsulated drug; furthermore, drug in nanoparticles could achieve the same effect of twice the concentration of unencapsulated drug [79]. Notably, as discussed above, only nanoparticles below a certain size (approximately 70 nm) are expected to be able to move through the brain tissue, but despite not being PEGylated or otherwise surface-modified, the PLGA nanoparticles in this study were able to move sufficiently through the interstitium with the aid of convective flow. Aside from use in delivering polymeric nanoparticles, CED has also been used to successfully demonstrate improved delivery of chemotherapeutic drug-loaded liposomes [80,81], inorganic nanoparticles for glioma imaging or therapy [82], drug/dendrimer conjugates [83], and even viruses for gene delivery [84]. Moreover, CED can be combined with other strategies mentioned above, such as nanoparticle surface modification, infusion with a hyper osmolar solution and ECM degradation to increase the size of the pores or channels through which nanoparticles can travel (Figure 3) [85].

It should be noted, however, that CED is not without risks. A Phase I/II clinical study using CED to enhance paclitaxel transport illustrated high efficacy in tumor reduction or slowed growth; however, complications also arose throughout the study in nine of 15 patients, a high rate that raises concern about translating this technology to the clinic [86]. Along with expected toxic side effects of the drug, there were instances of infection, accidental removal of the catheter, or loss of sufficient convective flux when infused fluid entered a cyst. As with other methods of local delivery, CED is an invasive procedure, and the amount of time that a patient is left exposed is intrinsically long due to the infusion process. Thus, CED has demonstrated promise in improving drug and nanoparticle transport through brain or tumor tissue, but safe ways of implementing the technology are still in development.

**Targeting brain tumor tissue over healthy brain tissue**

The strategies discussed thus far are generally all methods to cross the major physical barriers specific to brain or CNS delivery. However, as with other cancers, widespread toxic side effects, which may limit the tolerable dose of a drug, are an important problem that must be addressed in brain cancer treatment. Even if limited to the brain alone, minimal damage to healthy brain tissue must be a design goal for any therapy. Local delivery methods sidestep this issue to a large extent, although specificity for cancer cells at the site of the tumor could further ameliorate the problem. For systemic delivery, which has greater potential to reach metastatic cells or tumors that recur distant from the initial tumor site, the ability of a drug or drug carrier to affect the tumor preferentially to healthy cells would be a major advantage.

Nanoparticle–drug formulations are particularly exciting in this regard, as their design parameters can be modified and finely tuned to include one or several layers of delivery specificity. The EPR effect as described allows passive targeting of tumors by preferential extravasation and retention of nanoparticles within a certain size range. Numerous other targeting methods can be used individually or even in combination in the same nanoparticle, such as conjugation of or coating with a targeting ligand to enhance localization at the tumor; introducing degradable units or other properties to promote drug release near the tumor environment, capitalizing on microenvironmental factors such as decreased pH and
increased protease expression near tumors; and delivery of cargo that is preferentially or solely active in malignant cells. The last of these will be discussed further below in the context of particular therapeutics currently being studied.

Numerous examples exist of approaches to increase brain cancer specificity of nanomedicines. The folate receptor, as previously mentioned, has some specificity for tumors over healthy tissues, based primarily on the rapid rate of cell growth and division of cancer cells. Chitosan-based nanoparticles were conjugated with fluorescently labeled chlorotoxin, which binds specifically to gliomas and other tumors related in origin [87], and showed preferential accumulation in gliomas in mice [88,89]. Liposomes conjugated with IL-13 were also able to deliver doxorubicin specifically to glioma cell lines in vitro, based on the high expression of IL-13 receptor α2 on glioma cells but not on healthy brain tissue [90]. The CREKA peptide was previously shown to home toward tumor tissue and was coupled to 50-nm SPIO nanoparticles. When injected into a mouse, the particles not only targeted tumor vasculature but also induced clot formation, which in turn allowed binding of more circulating nanoparticles to the site [91]. Although this amplified targeting system was described in a breast cancer model, similar principles, using nanoparticles to incorporate many targeting functionalities, could potentially be applied to brain cancer as well.

Duan et al. developed pH-sensitive nanogels composed primarily of chitosan grafted to poly(N-isopropylacrylamide) and showed that release of the drug oridonin occurred preferentially in the pH range of 5.0–6.5 compared with 7.4 [92]. Because the pH surrounding tumors is typically <7.2, this strategy provides a possible method of targeting tumors [40]. In addition to changes in acidity, tumor cells often differ from healthy cells in that the former have higher intracellular concentrations of glutathione (GSH) [93], which can be used as a method to cause intracellular drug release preferentially in cancer cells. The high expression of GSH transporter on these cells also allows GSH to be used as a targeting ligand conjugated to such nanomedicines as the PEGylated liposomal doxorubicin for increased uptake [94], and this formulation has entered clinical trials [30].

Current & future drugs & experimental therapies in glioma

Nanoparticle-engineered cells as vehicles for glioma treatment

In recent years, some cell types have been found to migrate toward malignant GBs, providing an intriguing and promising option for improving efficacy and specificity of drug delivery to brain tumors. These include some adult stem cells such as neural stem cells (NSCs). It was found in 2000 that NSCs injected into a murine glioma model had high specificity for tumor tissue and the ability to spread throughout the tumor when NSCs were injected intratumorally or distant from the tumor, in either the ipsilateral or contralateral hemisphere, into the ventricles, or via tail-vein injection [95]. In highly invasive models, labeled NSCs were found to ‘track’ infiltrating glioma cells instead of staying merely in the tumor bulk. The authors of this study suggested that genetically engineered NSCs could be used to deliver anti-tumor agents, which they achieved in their study using retroviral gene delivery. Researchers have since discovered that other adult stem cells, such as bone marrow- (BM-MSCs) and adipose-derived mesenchymal stem cells, also display a high degree of tumor specificity in several glioma models, able to migrate across hemispheres and the BBB and toward the tumor [96]. MSCs are more readily available in large numbers than NSCs, potentially from the patient’s own bone marrow, blood or adipose tissue, and they can be expanded in culture.

MSCs have been proposed as a delivery vehicle to brain malignancies, either carrying drug- or imaging agent-loaded nanoparticles or engineered to secrete a therapeutic protein, with the advantage of having active migration mechanisms and not relying solely on diffusive
and convective transport through tissue as do nano-particles alone. For the latter, viruses – particles approximately 100 nm in diameter that have evolved for efficient gene transfer – have historically been used as gene delivery vectors because of their high efficiency. For instance, Nakamizo et al. also engineered their MSCs to overexpress interferon- θ(IFN- θ) using an adenoviral vector before injection, leading to improved survival over controls [96]. A number of groups have used retroviruses to transduce BM-MSCs [97] or adipose-derived mesenchymal stem cells [98] with the HSV-TK gene, which catalyzes a key step in converting the prodrug ganciclovir to a cytotoxic metabolite. These MSCs injected into the ventricles or lateral vein of tumor-bearing mice were shown to localize at the tumor and, upon injection of ganciclovir, reduce tumor growth by the bystander effect and improve survival.

Because nanoparticles are the appropriate size for cellular uptake while also being able to protect or tune the release of drugs encapsulated within, several groups have investigated the ability of MSCs to efficiently take up and carry cargo to brain tumors in the form of intact nanoparticles, which could then release a drug or perform a function at the tumor site. For example, oncolytic viruses can be loaded in vitro into BM-MSCs, which would carry the virus particles to the tumor site [99]. Although virus-mediated toxicity to the MSCs was a limiting factor, using MSCs as a delivery vehicle was superior in efficiency and specificity to delivery of oncolytic viruses alone. PLA nanoparticles loaded with fluorescent coumarin-6 could also be taken up by MSCs without affecting cell viability or phenotype; MSCs loaded with PLA and injected intratumorally in a mouse glioma model tended to stay near the tumor bulk as well as following behind infiltrating glioma cells [100]. SPIO nanoparticles loaded into labeled MSCs showed tumor homing when the loaded stem cells were injected systemically into a rat glioma model, providing a potential method for delivering nanoparticles to enhance MRI contrast [101]. Another group synthesized mesoporous silica nanoparticles (MSNs), loaded the particles with a mixture of different imaging contrast agents, and coated the loaded particles with hyaluronic acid for efficient uptake by BM-MSCs. These MSCs with loaded nanoparticles were injected systemically into mice and showed high specificity for the glioma site, as well as allowing in vivo imaging via near-infrared fluorescence, MRI and PET (Figure 4) [102]. Using adult stem cells as delivery vehicles is therefore a versatile and potentially powerful method for delivery of diagnostic or therapeutic nanoparticles to malignant gliomas.

**Nanoparticle formulations of conventional drugs**

As mentioned above, the drugs currently used for chemotherapy have made significant but small improvements in the prognosis of malignant glioma. Combinations of various therapies can further improve prognosis [103], but there is still a high need for more effective treatment. The current standard of care normally includes surgical resection with subsequent radiotherapy, often supplemented with adjuvant chemotherapy, including the locally implanted Gliadel BCNU wafer and the alkylating agent temozolomide.

Some compounds, including drugs used in the treatment of other cancers, are limited in use for glioma therapy because they cannot cross the BBB or are cleared too quickly for sufficient accumulation, they are not specific enough to prevent widespread toxicity, or tumor cells tend to be resistant to them. As a result, nanoparticles are being used as a method to reduce those problems with conventional drugs as well as to explore the potential use of other, more experimental therapeutic agents. For instance, the drug doxorubicin normally has poor to insignificant penetration of the BBB. By encapsulating doxorubicin into polysorbate 80-coated butylcyanocrylate nanoparticles, high uptake into the brain was seen in rats, with decreased systemic toxicity, increased survival rate and long-term remission in >20% of animals tested [104]. Efflux of drugs via the P-gp transporter, a mechanism of drug
resistance, was decreased by encapsulating paclitaxel in nanoparticles in polysorbate 80-coated nanoparticles [26]. The use of this nanoparticle system simultaneously decreased the toxicity due to Cremophor, which is usually needed for paclitaxel delivery, and also increased BBB penetration and decreased P-gp transporter activity due to the surfactant coating.

**Nanoparticles for gene therapy**

Nanoparticle-mediated DNA delivery was briefly discussed above in the context of engineering cells for glioma therapy, but biologics like DNA and multiple types of RNA are also under investigation as direct treatments for brain cancer. For nucleic acids, viruses have been the traditional method of delivery, as they have the advantages of high efficacy as well as the potential ability to cross various intra- and extra-cellular barriers. For example, clinical trials have demonstrated the efficacy of gene transfer treatment using an adenoviral vector. The adenovirus carried the HSV-TK gene and was injected into tumor margins following resection [105,106], exploiting the same TK/ganciclovir strategy described above and first proposed in the context of viral gene delivery in 1986 [107]. However, this and other studies have shown that replication-incompetent retroviruses have low efficacy in combating cancer [108]. Adenoviruses that do not express the E1B gene have been found to be selective for p53-deficient cells over the p53-expressing, healthy neural population, successfully targeting the tumor cells for viral infection and death in glioma patients [109]. In a study that integrated aspects of both of these approaches, Tai et al. used a retrovirus that selectively replicated in dividing cells to transduce glioma cells in a mouse with the cytosine deaminase suicide gene [110]. As a result, the transduced glioma cells were able to convert an intravenously injected prodrug, 5-fluorocytosine, to the toxic 5-fluorouracil, thereby causing local tumor death. In addition to conferring sensitivity to a drug or prodrug, viruses have also been used to deliver the p53 gene, a tumor suppressor that is commonly inactivated in gliomas [111]. Phase I clinical trials have used an adenoviral vector delivered intratumorally to transduce tumor cells with p53 [112,113], resulting in common but not prohibitively severe adverse side effects due to viral toxicity or immune response. Furthermore, preliminary results were suggestive of some anti-tumor effect.

It should be noted that, in one of the cases described above, four of 14 patients had an immune response to the adenovirus, and the frequency of seizures increased in two others. A Phase I/II trial using virus-producing cells following the HSV-TK/ganciclovir paradigm reported adverse side effects related to therapy in over 50% of patients [114], and worries about safety have limited the use of viruses in the clinic in general [115]. Aside from safety concerns, however, viruses are limited in their maximum cargo size and ease and consistency of manufacturing [116], leading to interest in the potential of using non-viral gene transfer agents. In a study that took place between 1999 and 2001 HSV-TK was delivered locally to recurrent GB patients using a liposomal formulation and showed increased necrosis to portions of the tumor after ganciclovir infusion, though not all of the patients appeared to have had sufficiently transfected tumor cell [117], and six of eight patients experienced adverse side effects, including fever and neurological deficits. Cationic liposomes carrying the IFN-β gene injected intratumorally into mouse models were also shown to be effective in reducing tumor growth and causing immune response in the tumor, and a pilot clinical trial in five malignant glioma patients based on these results showed >50% tumor reduction or stable disease in four of the patients [118]. In addition to liposomes, other materials have been used to make gene delivery nanoparticles for glioma therapy. PEG conjugated to PLA was used to encapsulate Apo2L/tumor necrosis factor-related apoptosis inducing ligand (TRAIL) plasmid into nano-particles <120 nm in diameter, which were then surface-conjugated to albumin. Because TRAIL is fairly specific for cancer cells over healthy tissue due to overexpression of TRAIL death receptors [119], most
resulting cell death was expected to be in the tumors. These nanoparticles were then injected intravenously and caused increased median survival time over controls [120].

Gene therapy also provides additional ways to introduce specificity into the system. As with the examples of TRAIL and p53 DNA, the protein product of the delivered gene should be designed to be active only in cancer cells. Moreover, the delivered construct can be under the control of a cancer-specific promoter, such as survivin [121] or PEG3 [122], ensuring that, should even healthy cells be transfected or transduced, the therapeutic protein would not be expressed. In addition to DNA delivery for gene overexpression, RNAi to knock down gene expression has in recent years been an active area of investigation. Cationic lipids and synthetic polymers have been used to form nanoparticle complexes with siRNA against a reporter gene as a proof-of-concept, showing effective gene knockdown in GB cells [123,124]. shRNA can also be expressed from a DNA plasmid, allowing the more extensively studied materials for stable DNA nanoparticles to be used for RNAi. Many potential molecular targets have been explored using nanoscale delivery [125]. A study using the cationic lipid Lipofectamine™ 2000 (Invitrogen, Paisley, UK) to encapsulate plasmids expressing siRNA against EGFR, a gene often mutated or overexpressed in malignant glioma [126], was able to decrease tumor volume a mouse glioma model up to fivefold (Figure 5) [127].

Inorganic nanoparticles: brain cancer imaging & treatment

Metallic nanoparticles have been studied in the laboratory as well as in the clinic. In one study, direct intratumoral injection of SPIO nanoparticles in a rat glioma model using RG-2 line tumors was followed by application of an alternating magnetic field that caused heat generation by the nanoparticles due to their increased Brownian motion [128]. The authors showed a 4.5-fold improvement in survival over untreated controls, although they did not compare their treatment directly with clinical gold standards like radiotherapy and/or conventional chemotherapy. Nevertheless, thermotherapy using this or similar methods using inorganic nanoparticles has high potential efficacy and has been studied by several groups. The use of super-paramagnetic particles as MRI contrast agents has been studied for over two decades [129], with many important physicochemical parameters for synthesis and imaging purposes already well understood. For example, the properties of these particles can be adjusted to target brain tumors using surface-conjugation of glioma-specific ligands [130], while other researchers use nanoparticles approximately 100 nm in diameter in order to take advantage of the EPR effect from disrupted BBB [35,131], providing MRI contrast of the tumor interstitium as well as providing a potential carrier for drug delivery.

Although inorganic nanoparticles are common in both research and the clinic, some question exists about the safety of their use [132], given that they are not biodegradable as are many nanoparticles currently in development. While several in vivo studies with metallic particles have shown no apparent toxicity, there is still concern about potential adverse side effects, particularly for inorganic particles that nonspecifically accumulate in non-target tissues or in healthy parts of the brain [133].

Brain tumor stem cells as a therapeutic target

Recurrence is an unfortunate certainty for most GB patients. One population of GB cells thought to be responsible in large part for this is the brain tumor stem cells (BTSCs). BTSCs within the bulk tumor have stem-like properties, including the ability to initiate and repopulate a tumor even most of it has been removed, and they have also been found to be refractory to many conventional anticancer treatments like BCNU [134,135]. They are therefore an attractive target for brain cancer therapy research, including experimental techniques that may be able to overcome the drawbacks of conventional chemotherapy and...
radio-therapy. Aside from inducing cell death in BTSCs, terminal differentiation of BTSCs could prevent surviving stem cells from being able to repopulate a tumor. For example, delivery of certain miRNAs such as miR-124 and miR-137 was found to cause terminal differentiation and cell death of murine BTSC mimics in vitro [136]. In this case, a lipid-based, commonly used commercial gene delivery agent, Lipofectamine formed nanocomplexes with miRNAs for successful in vitro knockdown. Gangemi et al. used shRNA-expressing plasmid in a retroviral vector for in vitro knockdown of SOX2, a transcription factor found in stem cells to be important in self-renewal, and found that this inhibited BTSC proliferation, self-renewal and tumor-initiating capacity [137]. Decreasing the stem-like properties of BTSCs in combination with the anti-tumor treatments described above may be an important and necessary aspect to add to current therapy regimens.

Future perspective

Brain cancer research and medical practice have advanced over the past decades, but progress, while significant, has been incremental and slow. The use of nanoparticles in this field has been fueled by a lack of current solutions to many of the barriers that impede further progress. Clinical translation of these technologies, in particular, has been slow, as few related studies have reached clinical trials and fewer still for applications in the brain [138]. Difficulties in translation of nanomedicine and its use in cancer therapy have been discussed in detail elsewhere [139].

Nanoparticles are, however, inherently well-suited for cancer therapy simply due to their small size and their highly tunable physical, chemical and biological properties. The potential therapeutic value of nanomedicine is huge as several functions for drug delivery and imaging can be simultaneously incorporated. Examples include multistep cancer targeting strategies and multimodal imaging agents as described above. With the additional consideration of using nano-particles to engineer targeting cells like MSCs or using targeting MSCs to deliver nanoparticles, even more potential treatment paradigms are possible. Nanotechnology can allow researchers to develop novel strategies for delivering drug cargos and imaging agents while also allowing the providing chemical flexibility to modify and functionalize nano particles, which may lead to a more comprehensive therapy for brain cancer.

Key Terms

<table>
<thead>
<tr>
<th>Term</th>
<th>Definition</th>
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<tbody>
<tr>
<td>Glioblastoma</td>
<td>Grade IV malignant glioma with high invasiveness and poor prognosis</td>
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<tr>
<td>Nanoparticle</td>
<td>Any particle in the size range of 1 nm–1 μm. The versatility of nanoparticle composition, structure, and carrying capacity makes them attractive as a potential therapeutic vehicle</td>
</tr>
<tr>
<td>Blood–brain barrier</td>
<td>Endothelial cells of brain vasculature prevent passage of most systemically delivered drugs from the circulation and into the brain</td>
</tr>
<tr>
<td>Enhanced permeability and retention effect</td>
<td>Disrupted, ‘leaky’ vasculature feeding tumors allows preferential accumulation of nanoparticles in the tumor microenvironment</td>
</tr>
<tr>
<td>Convection-enhanced delivery</td>
<td>Use of bulk fluid flow (convection) in addition to diffusion to promote movement of drugs and nanoparticles through the interstitium</td>
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</table>
**Brain tumor stem cell**

One of the population of cells in a tumor with the ability to initiate a new tumor. Brain tumor stem cells have been found to be resistant to many conventional therapies, which may contribute to tumor recurrence.

### References

Papers of special note have been highlighted as:

- □ of interest
- □□ of considerable interest

36. Sarin H, Kanevsky AS, Wu HT, et al. Physiologic upper limit of pore size in the blood-tumor barrier of malignant solid tumors. J Transl Med. 2009; 7:51. Although vascular abnormalities and the enhanced permeation and retention effect are important considerations in brain cancer therapy, this study emphasizes the strict size requirements of nanoparticles that can use the enhanced permeation and retention effect to cross the blood–brain barrier, even when compromised. [PubMed: 19549317]


delivery to the brain and also explains and overcomes physicochemical properties of nanoparticles, other than size, that restrict diffusion.


migratory abilities of injected stem cells and also their potential use as nanoengineered vehicles for drug delivery. [PubMed: 11070094]


*Ther Deliv.* Author manuscript; available in PMC 2014 April 01.


Executive summary

Shortcomings in current malignant brain cancer treatment

- Grade III (anaplastic astrocytoma) and IV (glioblastoma) gliomas have median survivals of 2–3 years and 15 months, respectively. Advances to the field in the past decades have improved prognosis only marginally.
- Experimental therapies such as nanotechnology may fill the gaps in the field.

Types of nanoparticles & their properties

- Biodegradable polymers (e.g., poly(ester)s, poly(amidoamine)s and chitosan) can form nanoparticles encapsulating small-molecule drugs, proteins, peptides or nucleic acids.
- Liposomes are commonly used but are cleared quickly by mononuclear phagocyte system cells. Modification with ‘stealth’ coatings such as polyethylene glycol can increase their half-life.
- Inorganic nanoparticles can be used as MRI contrast agents or as composite materials with multimodal function.
- Nanoparticles can protect therapeutic cargo from degradation or clearance. Their size and surface properties affect intracellular delivery efficacy (cellular uptake), biodistribution and clearance rates.

Blood–brain barrier

- Tight junctions between endothelial cells of capillaries in the brain prevent the exchange of many foreign materials, including systemically administered drugs.
- The blood–brain barrier (BBB) is disrupted at tumors, allowing passive nanoparticle accumulation via the enhanced permeability and retention effect.
- BBB disruption by hyperosmotic agents (e.g., mannitol) or vasodilators (e.g., bradykinin or histamine) increases the gap between endothelial cells; however, this does not distinguish between BBB at the tumor and the BBB of healthy brain tissue.
- Nanoparticles can be coated or conjugated to a ligand that penetrates the BBB.

Nanoparticle transport through brain interstitium

- For both systemic and local delivery, extracellular matrix prevents large particles from efficient diffusion through the brain. The maximum particle size can be increased by surface modification (e.g., PEGylation).
- Convection-enhanced delivery uses a hydrostatic pressure gradient to drive bulk flow in the tissue interstitium. Although this better distributes the nanoparticles, safety concerns have been seen in clinical trials.

Tumor-specific (targeted) delivery

- Ligands can target nanoparticles to fast-growing cells, which overexpress molecules like folate and transferrin receptors. Tumor-specific peptide sequences further increase specificity.
- Nanoparticle release can be targeted to the tumor environment (e.g., low pH).
The cargo being delivered can be specific to tumor tissue (e.g., transcriptional targeting with nucleic acids, proteins interacting with molecules overexpressed in tumors).

**Strategies under investigation**

- Common chemotherapy drugs such as doxorubicin have been encapsulated into nanoparticles that can penetrate the BBB.
- Viral and non-viral gene carriers can deliver a therapeutic to sensitize tumors to chemotherapy or to directly cause apoptosis.
- Neural stem cells and mesenchymal stem cells home toward tumors. Nanoparticles can be used to engineer these cells to express a therapeutic protein at the site of tumor.
- Nanoparticle-based therapies are being developed to target brain tumor stem cells to prevent recurrence.
Figure 1. Major barriers in nanoparticle delivery to malignant glioma
These barriers can be overcome by various particle modifications, such as ‘stealth’ surface coating, exploitation of the enhanced permeation and retention effect, or conjugation with BBB-penetrating or -binding molecules. (A) Clearance by immune cells; short half-life, (B) BBB, (C) insufficient diffusion through tissue, (D) non-specific delivery to healthy cells and (E) tumor invasion. BBB: Blood–brain barrier.
Figure 2. Targeted nanoparticle delivery to the brain
(A) Unmodified control nanoparticles demonstrated low accumulation in the brain compared with (B) RVG29-modified particles particles when injected intravenously into a mouse. Reproduced from [57] with permission from Elsevier.
Figure 3. Nanoparticle transport in the brain
Nanoparticles 30–40 nm in diameter were infused into the striatum of rats, either on their own in isotonic suspension (control), with hyperosmolar mannitol, or in isotonic suspension after pre-treatment with hyaluronidase or saline. The fluorescence image indicates the distribution of nanoparticles within 1 mm from the injection site. All three treatment conditions served to increase the effective pore size in the brain extracellular space during (mannitol) or prior to (enzyme, PBS) nanoparticle infusion, resulting in increased transport. PBS: Phosphate-buffered saline.
Reproduced from [85] with permission from Elsevier.
Figure 4. Mesoporous silica nanoparticles with multimodal imaging capacity demonstrated stem cell accumulation in tumors

MSNs were synthesized with two fluorescent dyes (FITC and ZW800, blue-green and near-infrared, respectively) and then loaded with $^{64}$Cu for MRI or Gd$^{3+}$ for positron emission tomography imaging. Bone marrow-derived MSCs were then loaded with the labeled mesoporous silica nanoparticles.  

(A) Diluting the number of cells demonstrated a corresponding decrease in ZW800 signal.  

(B) After tail-vein injection of loaded MSCs into a mouse orthotopic glioma model, positron emission tomography signal was higher in the tumor 24 h after injection than it was prior to injection.  

(C) MRI signal was higher in MSC/MSN-treated mice than in MSN-treated mice without MSCs.  

(D) Fluorescence microscopy on tissue sections also showed higher accumulation at the tumor site by MSC/MSNs than MSNs alone. HA: Hyaluronic acid; MSC: Mesenchymal stem cell; MSN: Mesoporous silica nanoparticles. Reproduced from [102] with permission from Elsevier.
Figure 5. Lipid-based nanoparticle containing plasmid coding for siRNA against EGF receptor demonstrated RNAi delivery as a genetic therapy for glioma. (A) Mice treated with these liposomes had tumors significantly smaller than those treated with liposomes containing scrambled siRNA sequence or left untreated. The histology of tumors in these mice also indicated (B) downregulated EGF receptor expression, (C) upregulated glial fibrillary acidic protein, indicating a change in phenotype, (D) increased apoptosis by TUNEL staining and (E) decreased proliferating cell nuclear antigen. Adapted with permission from [127].
Table 1

Strategies for nanoparticle delivery to the brain.

<table>
<thead>
<tr>
<th>Approaches to overcome</th>
<th>Examples</th>
<th>Ref.</th>
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<tbody>
<tr>
<td><strong>Short drug half-life</strong></td>
<td></td>
<td></td>
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<tr>
<td>Protection of therapeutic from degradation</td>
<td>Condensation of nucleic acid cargo with poly(beta-amino esters) or metalic nanoparticles</td>
<td>[14,124]</td>
</tr>
<tr>
<td>Increase circulation time</td>
<td>Surface coating with ‘stealth’ molecules such as PEG</td>
<td>[12,18]</td>
</tr>
<tr>
<td><strong>BBB</strong></td>
<td></td>
<td></td>
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<tr>
<td>Enhanced permeation and retention effect</td>
<td>Small (&lt;100 nm or &lt;20 nm) inorganic nanoparticles accumulate at the tumor site</td>
<td>[35,131]</td>
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<tr>
<td>Global BBB disruption</td>
<td>Use of hypertonic mannitol or vasoactive agents to increase BBB permeability</td>
<td>[29]</td>
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<tr>
<td>BBB-penetrating nanoparticles</td>
<td>Coating with ligands for the LDL receptor, such as apoA-I, apoE or angiopep-2</td>
<td>[43,44,45]</td>
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<td></td>
<td>Coating with surfactants that increase apolipoprotein deposition, such as poloxamer 188</td>
<td>[48,49]</td>
</tr>
<tr>
<td></td>
<td>Coating with cationized serum albumin</td>
<td>[54]</td>
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<td></td>
<td>Conjugation to peptides derived from BBB-penetrating viruses and toxins</td>
<td>[55,57,58]</td>
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<tr>
<td></td>
<td>Conjugation with ligands that bind BBB endothelial cells, such as transferrin and insulin receptors</td>
<td>[61,65,66]</td>
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<tr>
<td><strong>Insufficient transport through brain interstitium</strong></td>
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<tr>
<td>Increase nanoparticle diffusivity</td>
<td>Decrease nanoparticle size to &lt;70 nm</td>
<td>[75]</td>
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<td></td>
<td>Surface modification with a dense layer of PEG</td>
<td>[76]</td>
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<tr>
<td>Increase bulk fluid flow for nanoparticle convection</td>
<td>Convection-enhanced delivery of poly(lactic-co-glycolic acid), lipid-based, magnetic, dendrimer and virus nanoparticles</td>
<td>[79–84]</td>
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<tr>
<td><strong>Nonspecific delivery of drug to healthy tissue</strong></td>
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<tr>
<td>Local delivery</td>
<td>Direct injection of particles into tumor or tumor periphery</td>
<td>[79–84]</td>
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<tr>
<td>Surface modification with ligands targeting fast-growing cells</td>
<td>Conjugation of nanoparticles to ligands like transferrin or folate</td>
<td>[62,65–67]</td>
</tr>
<tr>
<td>Surface modification with ligands targeting cancer cells</td>
<td>Conjugation of nanoparticles to chlorotoxin, IL-13, CREKA peptide or glutathione</td>
<td>[87–91,94]</td>
</tr>
<tr>
<td>Preferential nanoparticle degradation or disassembly in tumor environment</td>
<td>Nanogels containing poly(N-isopropylacrylamide) for preferential release at lowered pH</td>
<td>[92]</td>
</tr>
<tr>
<td>Engineering of tumor-homing stem cells to act as delivery vehicles</td>
<td>Neutral stem cells and mesenchymal stem cells virally transduced to express cytokine deaminase or HSV-TK to sensitize to prodrugs 5-FC and ganciclovir</td>
<td>[95,97,98]</td>
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<td>Mesenchymal stem cells carry tumor-killing agent directly to brain tumor via transduction of IFN-Γor infection with oncolytic virus</td>
<td>[96,99]</td>
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<tr>
<td>Approaches to overcome</td>
<td>Examples</td>
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<tr>
<td></td>
<td>Mesenchymal stem cells carry nanoparticles loaded with imaging contrast agents</td>
<td>[100–102]</td>
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</table>

BBB: Blood–brain barrier.