Emerging Technologies for Delivery of Biotherapeutics and Gene Therapy Across the Blood–Brain Barrier

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Abstract
Antibody, immuno- and gene therapies developed for neurological indications face a delivery challenge posed by various anatomical and physiological barriers within the central nervous system (CNS); most notably, the blood–brain barrier (BBB). Emerging delivery technologies for biotherapeutics have focused on trans-cellular pathways across the BBB utilizing receptor-mediated transcytosis (RMT). ‘Traditionally’ targeted RMT receptors, transferrin receptor (TfR) and insulin receptor (IR), are ubiquitously expressed and pose numerous translational challenges during development, including species differences and safety risks. Recent advances in antibody engineering technologies and discoveries of RMT targets and BBB-crossing antibodies that are more BBB-selective have combined to create a new preclinical pipeline of BBB-crossing biotherapeutics with improved efficacy and safety. Novel BBB-selective RMT targets and carrier antibodies have exposed additional opportunities for re-targeting gene delivery vectors or nanocarriers for more efficient brain delivery. Emergence and refinement of core technologies of genetic engineering and editing as well as biomanufacturing of viral vectors and cell-derived products have de-risked the path to the development of systemic gene therapy approaches for the CNS. In particular, brain-tropic viral vectors and extracellular vesicles have recently expanded the repertoire of brain delivery strategies for biotherapeutics. Whereas protein biotherapeutics and bispecific antibodies enabled for BBB transcytosis are rapidly heading towards clinical trials, systemic gene therapy approaches for CNS will likely remain in research phase for the foreseeable future. The promise and limitations of these emerging cross-BBB delivery technologies are further discussed in this article.

Key Points

While receptor-mediated transcytosis (RMT) has been successfully exploited to deliver biotherapeutics across the blood–brain barrier (BBB), new RMT targets and advances in antibody engineering technologies are jointly contributing to an emerging pipeline of more specific and safer brain-penetrating biotherapeutics.

Certain viral delivery platforms capable of crossing the BBB are being refined and engineered to mitigate safety risks and enable systemic, targeted gene therapy for CNS diseases.

Cell-derived vesicles, such as exosomes, are emerging as ‘next generation’ biotherapeutic products capable of carrying versatile therapeutic payloads to target sites. Experimental demonstration of their ability to interact with and cross the BBB raises hope that the technology can be harnessed to devise novel strategies to deliver biotherapeutics across the BBB.

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Diseases of the CNS are among the most difficult to treat. Confounding factors include the complexity of brain physiology and pathology, as well as inadequate translational preclinical models to evaluate experimental therapeutics [1]. Despite significant progress in brain imaging surrogates for assessing therapeutic efficacy, more accessible brain-specific molecular biomarkers for early diagnosis and patient stratification for clinical trials remain sparse [1]. In addition, the delivery of therapeutics across the blood–brain barrier (BBB) remains one of the prime challenges in CNS drug development. The BBB is formed by specialized endothelial cells of brain microvessels and capillaries joined together by tight junctions that restrict paracellular transport of hydrophilic therapeutics and capillaries joined together by tight junctions that restrict paracellular transport of hydrophilic therapeutics > 500 Da [2]. Polarized efflux transporters further prevent brain access to many lipophilic synthetic molecules [2].

Biologics, pharmaceutical drug products manufactured in, extracted from, or semi-synthesized from biological sources, include vaccines, recombinant therapeutic proteins, gene therapy, and living cells or their products. Biologics are typically large and complex therapeutics, and their systemic delivery across the BBB was not considered feasible until recently. Alternative routes of delivery, including intraparenchymal pumps and intrathecal infusion, proved to be of limited value since the diffusion of these molecules within or into brain tissue, respectively, remained limited. Similarly, some clinically implemented medical procedures such as mannitol-induced osmotic BBB disruption [3] and focused ultrasound [4] for treatment of the brain have yet to show clinical benefits with biotherapeutics. Therefore, the development of novel transvascular brain delivery technologies remains critical for expanding the application of biologics, a highly successful therapeutic modality for cancer and inflammatory diseases, into diseases of the CNS.

Molecular Trojan horses are ligands or antibodies against the BBB receptors that transport essential growth factors and nutrients into the brain [5, 6]. The initial internalization and potential transport across the brain endothelium could occur via a non-specific, charge-mediated adsorptive endocytosis or via an energy-dependent and receptor-specific receptor-mediated endocytosis/transcytosis (RMT) [6]. Since the RMT process occurs via a specific receptor, it has potential for higher selectivity compared with the charge-mediated adsorptive process. The RMT process involves (i) engagement of the receptor on the luminal surface of brain endothelial cells (BEC) by the natural ligand or targeting moiety; (ii) endocytosis of the receptor-ligand complex into BEC; (iii) transcellular (transvascular) routing through complex endosomal sorting pathways, which may be receptor/cargo-specific; (iv) release of the cargo on the abluminal surface of BEC, and (v) subsequent receptor recycling to the luminal membrane of BEC [6] (Fig. 1a). The two most studied receptors undergoing RMT are transferrin receptor (TfR) and insulin receptor (IR) [6]. Antibodies against both of these receptors have been studied as molecular carriers to deliver attached therapeutic cargoes, including biologics, across the BBB. The enhanced and pharmacologically relevant brain delivery of biologics, including antibodies, growth factors, decoy receptors, and peptides has been demonstrated in pre-clinical models using various bioengineered antibodies against TfR [7–10]. Recent studies have focused on optimizing affinity of TfR antibodies [7, 10, 11] and used molecular modeling to establish the structure–function relationship of the receptor–antibody interactions (e.g., mono-valent vs bi-valent) that results in more efficient transcytosis [9]. These antibody engineering approaches resulted in significant improvements in TfR antibody pharmacokinetics [7, 10, 11], reduced lysosomal degradation within brain endothelial cells [11, 12], and enhanced efficiency of transcytosis and brain exposure [7, 9–11]. The molecular mechanisms that direct endosomal trafficking of receptor/antibody/cargo complexes towards the transcytosis pathway have been described in more detail recently [11–14]. Whereas TfR antibodies as molecular Trojan horses have still to enter clinical trials, a humanized mouse monoclonal antibody against IR fused to the enzyme iduronidase to enable its brain penetration in pediatric patients with severe Mucopolysaccharidosis Type I has been evaluated as safe and effective at improving biomarkers in an open-label, phase I–II clinical trial [15, 16].

However, the enthusiasm for TfR- and IR-targeting antibodies as BBB carriers has been dampened by translational gaps and safety concerns. Both receptors show species differences in abundance in brain endothelial cells [17], and antibodies raised against these receptors are typically species-selective, necessitating ‘surrogate’ molecules or humanization of the receptor in preclinical testing species. Additionally, translational gaps may be encountered due to neuroanatomical differences between pre-clinical animal models and humans, such as glia-neuron ratios [18]. Furthermore, safety concerns emerged during pre-clinical evaluation of TfR antibodies [19], stemming from the TfR enrichment in reticulocytes, the lungs and neurons [19, 20]. Thus, the quest continues for RMT receptors selectively expressed or highly enriched in BBB endothelial cells, which are expected to yield Trojan antibodies with better pharmacokinetic and safety profiles.

The recently demonstrated clinical benefit and subsequent FDA approval of intrathecal anti-sense oligonucleotide therapy (nusinersen, Spiraanza™) to correct a genetic mutation causing spinal muscular atrophy in pediatric patients [21], as
Fig. 1 A schematic depiction of the design and mechanisms of blood–brain barrier (BBB) transmigration of three emerging brain delivery technologies: BBB-crossing antibodies (a), brain-tropic adeno-associated viruses (b) and engineered extracellular vesicles (c, d).

a BBB-crossing antibodies are raised against a select number of BBB receptors that undergo receptor-mediated transcytosis (RMT) (Table 1). Therapeutic antibodies (TH-Ab) or other therapeutic cargoes, bio-engineered to incorporate BBB-crossing function, internalize via clathrin-coated vesicles and are trafficked through the early endosome pathway, including multivesicular bodies (MVB). They are released on the abluminal side of the BBB, as free antibodies or, in some cases, via exosomes, where therapeutic cargoes engage central targets (such as misfolded proteins or receptors on parenchymal cells) or replace missing molecules (for example growth factors or enzymes). The carrier receptor is recycled back to the luminal membranes via recycling endosomes to accept new cargoes from the circulatory compartment.

b Brain-tropic and non-brain-tropic AAVs, AAV9 and AAV2 respectively, recognize discrete receptors on brain endothelial cells, initiating internalization. Thereafter, distinct intracellular trafficking routes are utilized by AAV9 and AAV2, with the former undergoing active transcytosis across the BBB and the later peri-nuclear/nuclear localization and transduction of the endothelial cells. Following transcytosis across the BBB, AAV9 is taken up by and subsequently transduces parenchymal cells. AAV9 is also capable of spreading within the brain by anterograde, retrograde and trans-synaptic neuronal transport. Exosomes can be used as brain delivery vehicles in either their natural or in engineered forms. They can be loaded with various gene-based (siRNA, miRNA, DNA) and protein-based (antibodies, peptides) biotherapeutics. Specific targeting ligands are genetically engineered into the exosomal membranes in producing cells to achieve BBB crossing and/or drug delivery to target cells.

c Natural exosomes may internalize into brain endothelial cells by fusion with the plasma membrane and release their cargoes into endothelial cells. Exosomes expressing RMT ligands likely undergo clathrin-dependent endocytosis, trafficking and abluminal release. Transcytosing exosomes could target parenchymal cells or release their cargo into the brain extracellular space. AAV adeno-associated virus, MVB multivesicular body; TH Ab therapeutic antibody.
as well as promising results of the Ionis-Roche anti-sense oligonucleotide (RG6043) in phase I/IIa clinical trials for Huntington’s disease, leading to PRIME (PRiority MEdicines) designation by the European Medicines Agency, sparked a renewed promise that a broader application of gene therapy to correct devastating monogenetic neurological diseases, including lysosomal storage disease, ALS and Huntington’s disease, could be realized if appropriate brain delivery technologies were introduced in gene therapy designs. A major focus of recent gene therapy research has been in developing brain-tropic adeno-associated virus (AAV) vectors and understanding the mechanisms of their passage across the BBB [22]. While AAV-mediated gene therapy has great potential, significant obstacles remain to be surmounted in their development as therapeutics, including preclinical differences in pre-existing anti-AAV antibodies [23] and differential neuron/glial selectivity [24].

Finally, the field has witnessed an explosion of various nanoparticle delivery systems using diverse materials and designs to enable more efficient gene and large protein delivery across the BBB [25]. The inherent issues associated with this approach include complexity of composition and production, the absence of toxicology studies, and size limitations for nano-formulation transport across the BBB and within the brain extracellular space (35–60 nm) [26]. One promising novel approach relies on nanovesicles derived from cells, also known as exosomes, which could be enticed to package genes or other cargoes, as well as organ-targeting address molecules, by cell engineering and ex vivo bioprocessing.

The intensified research in BBB biology and pathology and advances in drug delivery and bio-technologies have jointly created critical mass in a field that has impeded CNS drug discovery for decades. Although the problem remains formidable, solutions have emerged that promise accelerated future development of BBB-crossing biotherapeutics for diseases of the CNS. Whereas advances in engineering and development of antibodies against IR and TfR have been the subject of numerous reviews in the past and recently, in this short overview we will focus primarily on emerging targets, technologies, and innovations in cross-BBB delivery arising from wide-scale genomics and screening studies.

2 Emerging Antibody Technologies for Brain Delivery Using Receptor-Mediated Transcytosis

The discovery of novel RMT receptor–antibody pairs has broadly followed two main workflows: (a) selection of potential RMT targets based on molecular (typically ‘omics’) analyses of the BBB and subsequent generation and evaluation of antibodies against these targets, and (b) ‘function-first’ screening of antibody libraries for antibody species that can transmigrate the BBB, followed by identification/deconvolution of the target(s) they engage. Both strategies have yielded initial proof of concept of enhanced brain delivery achieved with novel RMT targets and antibody modalities. The currently used RMT targets and antibody technologies and their characteristics are summarized in Table 1.

Using transcriptomic and proteomic analyses of mouse brain endothelial cells, Zuchero et al. [27] identified multiple highly abundant BBB-expressed proteins, including basigin, Glut1, and CD98hc. Brain penetration of the antibodies generated against these targets did not correlate with target expression abundance; however, antibodies against CD98hc showed a robust accumulation in the brain after systemic administration. When CD98hc antibodies were bioengineered as heterodimerized bi-specific antibodies with anti-BACE1 antibody, a strong pharmacodynamic response—a reduction in brain Aβ levels—was observed in transgenic animals after systemic administration [27] (Table 1). However, CD98hc antibodies accelerated systemic target-mediated clearance of the cargo antibody [27], due to the ubiquitous expression of CD98hc in peripheral tissues (Table 1). Antibodies against Glut1 and basigin showed a marginal increase in brain uptake, while antibodies against IR and Lrp1, also produced in this study, did not show improved brain penetration over that observed with a control antibody [27].

Similarly, using next-generation sequencing and proteomics analyses, we have selected insulin growth factor 1 receptor (IGF1R) as a potential target for RMT based on its abundance in isolated brain vessels compared with peripheral tissues. The panel of camelid single domain antibodies (VhHs) were raised against different epitopes of the extracellular domain of human IGF1R away from the IGF1 binding site, humanized, and evaluated for BBB crossing in models in vitro and in vivo [28] (Table 1). Selected IGF1R VhHs with species cross-reactivity demonstrated a saturable, energy-dependent transport across the human BBB model in vitro [29] and highly enhanced brain and CSF exposure in rats [29]. When used in monovalent fusion with antibody therapeutics, these VhHs had a minor impact on systemic pharmacokinetics, despite their high affinity. The transport of circulatory IGF1 across the BBB in vivo is driven by local neuronal activity [30]. Therefore, engagement of IGF1R by antibodies that do not interfere with IGF1 binding is a promising novel strategy to deliver biotherapeutic payloads across the BBB.

In separate studies using a ‘function-first’ screening approach to select BBB-crossing antibodies from high-diversity camelid VhH libraries, we identified the BBB-crossing antibody FC5 [31]. FC5 and its humanized variants have been genetically fused with various centrally
| RMT target                        | Target characteristics and function                                                                 | Antibodies against target                                                                 | Brain exposure and pharmacology                                                                 |
|----------------------------------|------------------------------------------------------------------------------------------------------|------------------------------------------------------------------------------------------|--------------------------------------------------------------------------------------------------|
| Transferrin receptor (TfR) [7–10, 19, 111] | Carrier protein for transferrin. Imports iron by internalizing the transferrin-iron complex through receptor-mediated endocytosis. TfR1 is ubiquitous; highly expressed in reticulocytes and lung, BBB and neurons. Soluble (circulatory) TfR levels used to assess iron status in patients. | Antibody formats include IgG, Fab, ScFv, single domain (VNAR) and F-star. Typically, not species cross-reactive. Biotechnologically engineered to bind the TfR in mono- or bi-valent format, and to have a range of affinities. Mono-valent TfR engagement and medium to low affinity range shown to improve BBB transcytosis. | Improved brain delivery and central pharmacodynamics of antibodies targeting amyloid beta and BACE1. High affinity TfR antibodies show fast systemic PK. Effector function-competent TfR antibodies show ‘on-target’ toxicity (reticulocyte lysis). |
| Insulin receptor (IR) [112–114] | Receptor tyrosine kinase; binding of insulin leads to phosphorylation of several intracellular substrates involved in metabolic actions of insulin, and cell growth and differentiation. Ubiquitously expressed. | Antibody formats include IgG, Fab, ScFv, single domain (VNAR) and F-star. Typically, not species cross-reactive. Biotechnologically engineered to bind the TfR in mono- or bi-valent format, and to have a range of affinities. Mono-valent TfR engagement and medium to low affinity range shown to improve BBB transcytosis. | Improved brain delivery and central pharmacodynamics of antibodies targeting amyloid beta and BACE1. High affinity TfR antibodies show fast systemic PK. Effector function-competent TfR antibodies show ‘on-target’ toxicity (reticulocyte lysis). |
| CD98hc [27]                     | The heavy chain of a heterodimer, covalently bound through di-sulfide bonds to one of several possible light chains; required for function of light chain amino-acid transporters (large neutral amino acids). When associated with SLC7A5/LAT1 also involved in the transport of L-DOPA. Ubiquitously expressed. | Antibodies generated using the murine extracellular domain to immunize mice, rats, or hamsters. Used as chimeric antibody on human Fc in either bi-valent or monovalent format. | Enhanced brain uptake in non-human primates, estimated at 1% ID. Delivery of enzymes for LSD demonstrated in NHPs; fast systemic PK. Safety and proof of concept clinical studies ongoing. |
| IGFI R [28]                     | Receptor tyrosine kinase; binds insulin-like growth factor 1 (IGF1) with high affinity and IGF2 and insulin (INS) with a lower affinity. The activated IGFI R is involved in cell growth and survival control. Expresses in periphery; however lower liver expression. Expressed in subsets of neuronal cells. | A panel of humanized camelid single-domain antibodies (VHHs) recognizing different epitopes on the α-subunit of the IGFI R, away from IGF1-binding site. Selective for IGFI R; some antibodies in the panel are species cross-reactive and pH sensitive. Binding affinities in low nanomolar range. Used in fusion with Fc fragment or with full antibodies (mono- or bi-valent). | Tolerates fusion on N- or C-terminus. When fused to Fc, shows up to 10-fold improved brain exposure compared to control antibodies. Mild target-mediated clearance in periphery. |
| TMEM30A (cdc50a) [31–33, 115] | Accessory (β subunit) of a P4-ATPase flipase which catalyzes transport of aminophospholipids from the outer to inner leaflet of various membranes. Implicated in vesicle formation and uptake of lipid signaling molecules. Ubiquitously expressed; enriched in brain vessels. | Humanized VHH isolated by ‘function first’ cell-based panning. Broadly species cross-reactive; apparent binding affinity 50–100 nM. Used in fusion with Fc fragment or with full bi-valent antibodies. | Tolerates fusion on C-terminus only. Improves brain delivery of full antibodies (mGluR1) and neuropeptides fused via an Fc fragment. Produces measurable cargo-mediated pharmacological responses elicited by central target engagement. |

BBB blood–brain barrier, RMT receptor-mediated transcytosis
acting payloads, including neuropeptides [32] and full monoclonal antibodies [33] (Table 1). FC5 had a minimal effect on circulatory pharmacokinetics of monoclonal antibodies, while increasing both their brain and CSF exposure in rats [33]. Target engagement and pharmacological effects of an FC5-delivered cargo antibody, acting as an antagonist of the G-protein-coupled metabotropic glutamate receptor 1 (mGluR1), have been observed in a rat model of thermal hyperalgesia; the bi-specific FC5-mGluR1 antibody co-localized with thalamic neurons involved in nociceptive responses through mGluR1 [33]. Deconvolution of the BBB receptor engaged by FC5 indicated a role for TMEM30A (cdc50A), a β subunit of phospholipid flippases that trigger endocytic vesicle formation by catalyzing transport of aminophospholipids from the outer to the inner leaflet of various membranes [34].

The above studies have provided a new generation of RMT targets and antibodies suitable for development of BBB delivery strategies for a wide range of biotherapeutics. Advances in the antibody engineering field have expanded formats of BBB-transporting antibodies from a full IgG, used to target IR, to antibody fragments (Fab, and ScFvs) and small monomeric single-domain antibodies of only 15 kDa, and, more recently, to antigen-binding sites incorporated into the Fc region of antibodies (Fcab™).

The engineering strategies to modulate the function of Fc fragments have also come into sharp focus. For example, an effector function of Fc may be required for therapeutic efficacy of the antibody in the brain—such is the case for amyloid β-targeting antibodies, where mobilization of the innate immune system and microglia via effector-competent Fc accelerates amyloid degradation and clearance; yet intact effector function may lead to systemic side effects when BBB carrier antibody binds peripheral targets. The example of the later is acute reticulocyte lysis observed with effector-competent Tfr-BACE-1 bi-specific antibodies [19]. However, mono-valent anti-Tfr Fab linked to a C-terminus of the Aβ-antibody could not trigger effector function systemically due to steric hindrance, while the bispecific antibody retained full effector function activity when engaging its brain target [33]. In addition, treatments of chronic diseases that require a constant presence of the therapeutic in the brain may benefit from an extended circulation half-life that can be achieved by Fc engineering.

Using these innovations and highly modular technologies in antibody engineering, various types of bi-specific or multi-specific antibodies that incorporate a BBB-crossing function and CNS-targeting function(s) have been engineered and expressed, as reviewed in [35], and are steadily progressing through pre-clinical development towards clinical trials.

3 Gene Therapy: Brain Tropic Adenoviral Vectors

The BBB presents a uniquely difficult challenge for the delivery of gene therapy to the CNS. Additionally, major challenges for viral gene therapy include peripheral and CNS off-target effects due to broad viral tropism, toxicity related to high gene transfer efficiency, insertional genotoxicity and anti-vector immune responses. Strategies are being developed to decrease off-target effects, including the incorporation of tissue-specific promoters for payloads [36], miRNA that suppress ectopic expression [37], as well as investigation of the role of adsorptive endocytosis in mediating brain endothelial cell transcytosis [38, 39].

Several approaches can be taken to manipulate the genome, including (i) silencing mutant alleles, (ii) replacement of dysfunctional mutated genes, (iii) introduction of a disease-modifying gene, and (iv) the correction of genetic mutations using gene editing methods. Compared with non-viral vectors, including the use of liposomes, exosomes, and polymeric nanoparticles [40, 41], viral vectors have proven more efficient in delivering gene therapy in vivo, where the goal is to transduce the maximal number of target cells [42].

Among various viral delivery systems summarized in Table 2, only select classes are suitable for CNS applications. For example, retroviruses are not suitable for use in neurological disorders because of their inability to transduce non-dividing cells. In comparison, adenoviruses have the advantages of the absence of genomic integration [43] and a very large carrying capacity, important for those diseases that require the delivery of large payloads, such as Huntington’s disease or Duchenne muscular dystrophy. However, adenoviruses are known to elicit a strong innate immune response, and severe cytotoxicity and inflammation were observed with high doses in a clinical trial [44].

AAV is a naturally replication-defective, non-pathogenic, non-enveloped single-stranded DNA parvovirus that has gained traction as a vector for CNS-targeted gene therapy [45]. Cellular transduction by AAV involves a series of events, including binding of a cell-surface receptor, internalization through endocytic uptake, endosomal/lysosomal escape, nuclear entry, capsid uncoating, and genome release [46, 47]. Events occurring during endosomal processing are indispensable to AAV transduction. Following release of the genome in the nucleus, second-strand synthesis and transcription ensue [48]. AAV exists primarily as an episomal vector that does not disrupt the host genome and is not burdened with the risk of activating oncogenes [49]. While persistent AAV expression can be achieved in non-dividing cells in the CNS, obtaining persistent expression in dividing cells requires additional modifications to the viral genome to enable chromosomal integration or episomal replication.
Table 2  Viral vector characteristics

| Viral vector                  | Particle size (nm) | Cargo          | Capacity (kb) | BBB-crossing | Axonal transport | Post-mitotic infection | Chromosomal integration | Immunogenic |
|-------------------------------|-------------------|----------------|---------------|--------------|------------------|------------------------|-------------------------|-------------|
| Adenovirus                    | 70–120            | dsDNA          | 8–10          | No           | No               | No                     | No                      | Strong      |
| Helper-dependent adenovirus   | 70–120            | dsDNA          | Up to 36      | No           | No               | No                     | Yes                     | Strong      |
| MoMLV retrovirus              | 100               | RNA            | 7–8           | No           | No               | No                     | Yes                     | Low         |
| Lentivirus                    | 100               | RNA            | 7–9           | No           | No               | Yes                    | Yes                     | Low         |
| Herpes simplex virus          | 120–300           | dsDNA          | 30–50         | BBB damage   | Yes              | Yes                    | Yes                     | Low         |
| Adeno-associated virus        | 20–25             | ssDNA          | 4.8           | Yes          | Yes              | Yes                    | Yes                     | Low         |

**BBB** blood-brain barrier, **dsDNA** double-stranded deoxyribonucleic acid, **ssDNA** single-stranded deoxyribonucleic acid, **RNA** ribonucleic acid

[50]. Although AAV has a small packaging capacity (4.8 kilobase pairs of nucleic acid), its ability to transduce mitotic and post-mitotic cells, tropism for neurons and glia, absence of pathogenicity or cytotoxicity, and low immunogenicity [51] are distinct benefits for CNS gene therapy [42]. A common method for optimizing gene expression is the use of self-complimentary AAV (scAAV) genomes, which can expedite expression and increase transduction efficiency but comes at a cost of a > 50% reduction in packaging capacity [52]. Additionally, the methods used for viral expression and purification have been shown to alter transduction patterns and efficiency [53], indicating a need for optimization and standardization in order to improve the reliability and predictability of AVV therapeutics.

### 3.1 Brain Tropism of Adeno-Associated Virus (AAV) Serotypes

AAV is a highly diverse group of viruses that include nine major serotypes (AAV1–9) and a variety of novel hybrid serotypes that are designed to alter viral tropism [54]. Typically, AAV utilizes cell-surface proteoglycans as a primary receptor along with a distinct co-receptor, and a given serotype can interact with multiple primary receptors. Thus, tissue tropism is dictated by the expression of the primary and secondary AAV receptors. A comprehensive study characterizing the in vivo biodistribution of AAV serotypes delivered via tail vein injection [55] demonstrated that the distribution and kinetics of expression varied widely among serotypes, with AAV9 found to produce the most robust tissue expression and, unlike other serotypes, was capable of producing expression in the brain.

Studies utilizing intraparenchymal injection of AAV have revealed that all serotypes are capable of transducing expression in brain cells of rodents [56, 57], non-human primates [58], and dogs [59]. However, AAV4 appeared to primarily target ependymal cells [60], AAV8 had the greatest tropism for astrocytes, microglia, and oligodendrocytes, and AAV9 had the greatest tropism for neurons [56]. Additionally, AAV9 produced the greatest distribution in brain tissue, likely related to its ability to undergo both anterograde and retrograde axonal transport as well as trans-synaptic transport (Fig. 1b) [56, 61]. AAV serotypes also display distinct transduction characteristics when administered in specific brain regions [56, 62]. While extensive variability between brains injected with AAV is reported [62], general trends are observed. Aschauer et al. [56] reported rankings of serotypes in three brain regions, finding that AAV9, AAV8, and AAV-5 produced high levels of expression, whereas AAVs consistently produced the lowest levels of expression in the striatum, hippocampus, and auditory cortex of adult mice.

Studies in rodents and non-human primates have conclusively shown that AAV9 is capable of crossing the BBB and transducing parenchymal cells, primarily neurons and glia [22, 24, 56, 63, 64], following systemic administration. Additionally, AAVrh8 and AAVrh10 have been demonstrated to produce robust expression in the CNS following intravascular administration without disruption of the BBB [63, 65, 66]. It should be noted that peripheral administration of AAV9 is also known to produce high levels of transduction in the liver and spleen [24].

AAV9 tropism is affected by both the route of administration and the age of the recipient animal. The brain regions to which AAV9 can spread following intraparenchymal injection are contingent upon the location of the injection, where spreading is dictated by the efferent and afferent projections in the injection region [67]. It has been consistently reported that intravascular administration of AAV9 transduces neurons and astrocytes in the brain and motor neurons in the spinal cord in animals (rodents and nonhuman primates); however, despite somewhat contradictory reports, the efficiency of neuronal transductions appears to decline with age [48].

Recently, progress has been made in understanding the mechanism by which AAV9 crosses the BBB. The observation that mannitol does not enhance brain delivery of AAV9, in contrast to AAV2, indicated that the mechanism of AAV9 BBB transport is via a transcellular pathway [24]. AAV particles can undergo cellular uptake through a
variety of processes, including macropinocytosis, phagocytosis, and clathrin- or caveolae-mediated endocytosis, with various serotypes exhibiting distinct transduction and entry profiles [47, 68] (Fig. 1b). For example, two distinct endocytic mechanisms mediate AAV2 uptake, namely CLIC/GEEC and dynamin-dependent endocytosis, with only the former resulting in transduction [47]. In a recent study by Merkel et al. [38], the transit of AAV9 particles across the human brain microvascular endothelial cell barrier required an active transport process whereas the low level transit of AAV2 involved an energy-independent paracellular diffusion process. While both AAV2 and AAV9 were taken up by endothelial cells, AAV2 was present in endosomal structures and accumulated in the perinuclear/nuclear region, and AAV9 was almost exclusively distributed to the basolateral surface of the endothelial cells, or in funnel-like structures within the endothelial cells. The differences in the intracellular trafficking of AAV2 and AAV9 are thought to be a consequence of the differences in their receptors/co-receptors, which presumably direct the particles to different endocytic mechanisms [47].

It is conceptually possible to engineer AAV capsids that interact with receptor systems that are restricted to brain endothelial cells and are capable of triggering receptor-mediated transcytosis. This principle was demonstrated using a directed evolution strategy, where AAV capsid shuffling was employed to generate vectors that selectively cross seizure-compromised BBB and exhibit a favorable biodistribution in rats [69]. Others [70–72] have utilized in vivo phage-display bio-panning to identify peptides that exhibit BBB specificity. Insertion of BBB-specific peptide motifs into the VP3 region of the AAV capsid enhanced the BBB tropism of AAV. Furthermore, Chen et al. [71] were able to create peptide-modified AAV with normal and disease-specific BBB tropism. Further refining this strategy to exploit known BBB transcytosis receptors should similarly enhance the specificity of BBB targeting. Consideration must also be given to the potential for adverse effects, such as increased immunogenicity or decreased transduction efficiency and BBB integrity, resulting from the introduction of non-natural sequence modifications. However, it has been demonstrated that directed evolution can be used to overcome immunogenicity [73], further indicating that the AAV capsid is amenable to engineering of beneficial functionality. Enhanced transduction of permissive and non-permissive cells following incorporation of cell-permeable peptides into the AAV capsid has been reported [74, 75]. However, this approach is not suitable for systemic administration as it currently lacks tissue specificity and would likely result in increased off-target transduction of peripheral tissues at the expense of RMT-mediated brain delivery.

### 3.2 Cargoes Compatible with AAV

While in vivo gene therapy based on viral vectors is thought to be best suited to gene addition/replacement [45, 76, 77], AAV can also be used to deliver gene-modifying oligonucleotides, such as short hairpin RNA (shRNA), microRNA, and antisense oligonucleotides (ASOs). The approval of nusinersen in 2017 demonstrates the validity of using ASOs to regulate exon splicing in diseases such as spinal muscular atrophy, Leber congenital amaurosis, muscular dystrophy, β-thalassemia, and spinocerebellar ataxia type 3. Similarly, correcting diseases by reducing aberrant overexpression can be achieved using ASOs, shRNA, and miRNA. AAV-mediated delivery of therapeutic ASOs may represent a means to produce sustained drug delivery at a more reasonable cost. AAV delivery of shRNA and miRNA has been successfully used to decrease in vivo expression of DNM2 (myotubular myopathy) [78] and mutant huntingtin [79] in mice. SOD1 was successfully targeted for reduction with an AAV-delivered exon skipping ASO in a murine model of amyotrophic lateral sclerosis [80]. These studies indicate that AAV-mediated delivery of gene-regulating oligonucleotides represents a viable strategy for delivering cost-effective DNA therapies for a wide spectrum of CNS diseases.

### 4 Cell-Produced Delivery Vehicles: Promise of Extracellular Vesicles

Exosomes have recently emerged as novel lipid nanocarriers for the delivery of small molecule drugs, protein- [81, 82], and gene-based biotherapeutics [83] (Fig. 1c–d). Exosomes are a sub-population of specialized membranous extracellular vesicles (EVs) derived from endocytic compartments, 30–100 nm in size, that are actively secreted by almost all cell types [84]. They are formed in multi-vesicular bodies and released into the extracellular space by fusion with the plasma membrane [85]. Exosomes express a ubiquitous set of pan-exosome proteins such as CD9, CD81, CD63, Alix, TSG101, HSP70, and flotillin-1; they also express cell-type-specific proteins and carry cell-specific miRNA cargoes that affect their function. Exosomes produced by HEK293T cells have been characterized and a comprehensive database of protein, mRNA, and miRNA profiles has been generated [86]. Exosomes derived from human brain endothelial cells have been shown to contain distinct, possibly tissue-specific biomarkers, as well as known RMT receptors including TIR, IR, LRP, and TMEM30A [87].

Exosomes participate in inter-cellular communication and transfer of cellular contents to adjacent cells or distant organs [88], which could subsequently modify target-cell
confocal microscopy examination demonstrated that inter-
shown to involve an energy-dependent endocytosis process; 
healthy BBB. The uptake of exosomes by HBMEC was 
shown to only cross TNFα-treated HBMECs but not the 
in vitro transwell model using HBMEC. Exosomes were 
across the healthy and inflamed BBB [93] was studied in 
HEK293T cells engineered to express luciferase (exo-luc) 
understood. The trafficking of exosomes derived from 
exosomes interact with brain endothelial cells are not well 
properties of exosomes have been exploited for development 
of delivery strategies for exogenous gene-based cargoes.

Brain- and BBB-tropism of exosomes derived from vari-
cious cellular sources has been studied in model organisms, 
such as zebrafish [91, 92]. Interestingly, among exosomes 
isolated from glioblastoma (U-87MG, A-172), neuroecto-
dermal (PFSK-1), and mouse brain endothelial (bEnd.3) 
cells, only brain endothelial cell-derived exosomes crossed 
the BBB and delivered encapsulated payloads, suggesting 
potential homo-tropism for the tissue of origin [92].

The mechanisms by which homo- or heterotypic 
exosomes interact with brain endothelial cells are not well 
understood. The trafficking of exosomes derived from 
HEK293T cells engineered to express luciferase (exo-luc) 
across the healthy and inflamed BBB [93] was studied in 
an in vitro transwell model using HBMEC. Exosomes were 
shown to only cross TNFα-treated HBMECs but not the 
healthy BBB. The uptake of exosomes by HBMEC was 
shown to involve an energy-dependent endocytosis process; 
confocal microscopy examination demonstrated that internal-
ized exosomes colocalized with transferrin-Texas Red, 
an early endosome marker, and also with cholera toxin B, 
a late endosomal marker. Although luciferase activity was 
detected in the abluminal chamber, it remained uncertain if it 
was associated with intact exosomes. Other studies have 
shown that exosomes can be internalized into cells by dif-
ferent mechanisms, including plasma membrane fusion, 
cathrin-dependent endocytosis and clathrin-independent 
pathways such as phagocytosis, macropinocytosis, caveo-
lin-mediated uptake, and lipid raft-mediated internalization 
[94–98].

An attractive strategy to achieve BBB crossing and cell-
type-specific delivery is to genetically engineer targeting 
ligands known to be involved in RMT-mediated transcy-
tosis (some shown in Table 1) into exosomal membrane 
proteins [83] (Fig. 1c). The preclinical proof-of-principle 
of this approach was demonstrated with exosomes derived 
from dendritic cells engineered to express lysosomal-asso-
ciated membrane protein 2b (Lamp2b) fused to the neuron-
specific rabies virus glycoprotein (RVG), and loaded with 
siRNA against BACE1 by electroporation [99]. Systemically 
injected RVG-siRNA exosomes silenced the mRNA and pro-
tein levels of BACE1 in neuronal cells, resulting in reduced 
brain amyloid levels in transgenic mice. The same strategy 
was also applied to knockdown the expression of α-synuclein 
in the mouse brain following intravenous administration of 
RVG-exosomes loaded with siRNA [100]. Although these 
preclinical studies are encouraging, many hurdles remain 
in the clinical translation of these findings [101].

4.1 Exosome-Associated Adeno-Associated Viruses (Exo-AAV)

Encapsulation within EVs has shown promise for increas-
ing AAV delivery to difficult targets. During the standard 
AAV vector production, a portion of AAV has been shown to 
associate with HEK293T producer cell exosomes (exo-AAV) 
[102], with AAV capsids present both on the surface and 
encapsulated inside exosomes [103]. Exosome encapsulation 
shields AAV from neutralizing antibodies, while enhancing 
AAV9 transduction in vitro and in vivo [104, 105].

In a recent study [105], both exo-AAV8 and exo-AAV9 
encoding green fluorescent protein (GFP) targeted neurons 
and astrocytes with higher transduction efficiency compared 
with conventional AAV8 or AAV9 after intravenous admin-
istration in Balb/c mice. Exo-AAV9 and exo-AAV8 had 9.3-
fold and 4.3-fold increased transport, respectively, across an 
in vitro BBB model using bEnd.3 cells, compared with their 
free viral counterparts. Similarly, exo-AAV2-GFP [106] and 
exo-AAV1 or exo-AAV9 vectors [107] efficiently targeted 
and delivered transgenes across retinal and cochlear brain 
barriers, respectively.

Exosomes can be isolated in large quantities from cells 
ex vivo and engineered to package specific cargoes into 
secreted macrovesicles through bioprocess scale-up, simi-
lar to that of virus-like particles (VLPs). The HEK293 cell 
platform has been widely used for the production of AAVs 
and therapeutic proteins that have been approved by the FDA 
[108]. However, there are no cell sources or manufacturing 
processes that have been FDA-approved for clinical-grade 
exosome production [109]. The International Society for 
Extracellular Vesicles (ISEV) has recently summarized the 
criteria that need to be met by exosome-based therapeutics to 
turn promising preclinical compounds into drug candidates 
[110]. Exosomes as ‘therapeutics’ are being tested in a few 
clinical trials currently; notably, benefits of plant exosomes 
or exosomes derived from allogeneic mesenchymal stromal 
cells are being evaluated in acute ischemic stroke, diabetes 
meilitus, and several cancer indications.

5 A Glimpse into the Future of CNS 
Biotherapeutics

With antibody technologies coming of age, supported by 
unprecedented advancements and precision in antibody 
engineering and biomanufacturing, and progress being 
made in deciphering targets and mechanisms of transcyto-
sis across the BBB, it is easy to imagine a future in which 
the development of all CNS biotherapeutics will be closely 
tied with the concurrent development and deployment of the 
tailored BBB-delivery strategy. The bi-specific antibodies 
and BBB carrier-fusion proteins targeting a variety of CNS
diseases, most notably those characterized by accumulation of misfolded proteins, are advancing through pre-clinical stages and some will be entering clinical trials as early as 2019. With this burgeoning pipeline, we can expect a proof of concept of the receptor-mediated transcytosis approach in humans in the near future.

Following closely behind is the wave of gene therapies for CNS diseases. Combining the use of gene editing tools and neuro-tropic viral vectors has placed gene therapy in a position to make breakthrough advances in the treatment of currently incurable monogenic brain diseases. Given initial successes with intrathecal administration of ASOs therapies, it is likely that this route of administration will dominate subsequent clinical trials of gene therapy. However, the progress made in engineering viral vectors with brain tropism and brain delivery technologies for RNA therapeutics will eventually shift the preference towards systemic administration, the only route that can successfully address the access of these therapies to deep brain structures in humans.

As the field of biotechnology embraces the complexity of future therapeutics, engineered cell-derived targeted nanovesicles may complement antibody and gene therapy strategies to enhance the versatility of delivery options. Emerging biotechnology companies in this field, such as Evox Therapeutics (https://www.evoxtherapeutics.com/), are engineering the natural delivery capabilities of extracellular vesicles to develop an entirely novel class of biotherapeutics, notably targeting CNS indications. Lastly, RMT-targeted exo-AAVs are evolving as particularly promising novel gene therapy delivery technology, since they possess beneficial features of both exosomes, such as low immunogenicity, and viral vectors, such as robust gene delivery.

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Compliance with Ethical Standards

Conflict of interest All authors are employees of the Government of Canada. None have any existing conflict of interest with regard to this manuscript.

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