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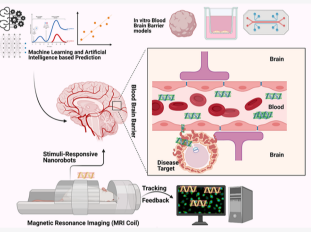
Review

Emerging Application of Nanorobotics and Artificial Intelligence To Cross the BBB: Advances in Design, Controlled Maneuvering, and Targeting of the Barriers

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ABSTRACT: The blood-brain barrier (BBB) is a prime focus for clinicians to maintain the homeostatic function in health and deliver the theranostics in brain cancer and number of neurological diseases. The structural hierarchy and in situ biochemical signaling of BBB neurovascular unit have been primary targets to recapitulate into the in vitro modules. The microengineered perfusion systems and development in 3D cellular and organoid culture have given a major thrust to BBB research for neuropharmacology. In this review, we focus on revisiting the nanoparticles based bimolecular engineering to enable them to maneuver, control, target, and deliver the theranostic payloads across cellular BBB as nanorobots or nanobots. Subsequently we provide a brief outline of specific case studies addressing the payload delivery in brain tumor and neurological disorders (e.g., Alzheimer's disease, Parkinson's disease, multiple sclerosis, etc.). In addition, we also address the opportunities and challenges across the nanorobots' development and design. Finally, we address how



computationally powered machine learning (ML) tools and artificial intelligence (AI) can be partnered with robotics to predict and design the next generation nanorobots to interact and deliver across the BBB without causing damage, toxicity, or malfunctions. The content of this review could be references to multidisciplinary science to clinicians, roboticists, chemists, and bioengineers involved in cutting-edge pharmaceutical design and BBB research.

KEYWORDS: Blood-brain barrier, nanorobots, transcytosis, machine learning and artificial intelligence, bioengineering, nanoparticles

1. INTRODUCTION

With the increase in neurological disorders and the demand for new drug development, a focus on expedite brain research can be seen in recent times. Alarming attention to develop novel neuropharmaceutic raised the urgency to develop in vitro models, mimicking in vivo like blood—brain barrier (BBB) structure/function relationship. Due to the complexity and poor accessibility, BBB models act as an effective alternate tool for brain research. Use of animal models is considered as the exemplar of drug testing as the BBB complexity can be easily recapitulated. This is contemplated to be highly advantageous considering the ease of testing of pharmaceutical interventions from the cellular to the systemic level. However, testing of drugs with in vivo models for BBB is highly tedious, immensely costly, and time-consuming. Further, there is a huge difference in drug interactions between animal model and human trials.^{1,2} Hence, researchers have started focusing on developing in vitro BBB models or BBB-on-chip devices to overcome these limitations of new drug development. The need for in vitro BBB models is indispensable as they also contribute in identifying the specific physiological and pathological mechanisms under diseased conditions, thereby uncovering the route to effective drug discovery. The challenges in modeling BBB by incorporating the key in vivo properties stem from the complex subtleties of BBB structure and the interactions with

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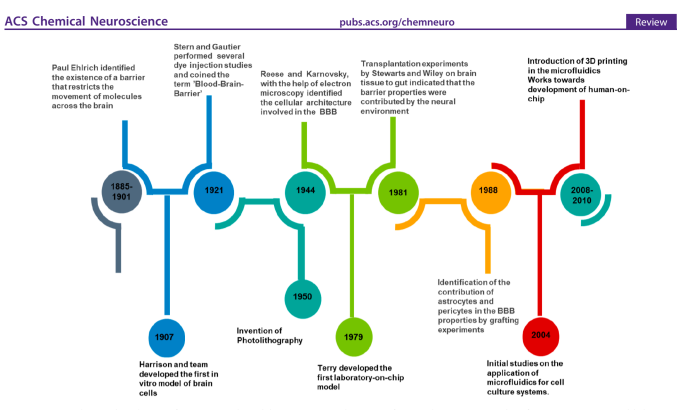


Figure 1. Time line and evolution of experimental model in BBB research ranging from early attempts to identify BBB existence, establishing its relation with CNS to the most advanced perfusion based models.

neighboring cells. Currently, complete mimicking of the entire brain structure and function in a laboratory engineered design is unavailable, due to gaps in the knowledge on complexity and diversity of brain function during healthy and diseased states. Further, limited techniques and their workability have provided scope for recapitulation of specific parts of brain like nonleaky and tight BBB.

Advent of nanocarrier systems for drug delivery has been a crowning discovery in the field of brain research as they are said to have the capabilities for conquering BBB defenses based on their surface properties. Nanoparticles, when combined with other elements like polymer, have shown ability to effectively cross BBB and deliver drugs to CNS.³ The many advantages of using nanoscale materials in brain research have been identified by several researchers,^{4,5} and many are still probing the potential of these materials.⁶ Nanorobotics represent a relatively recent advancement in this regard as they are engineered specifically with sensing, decision making, and actuation properties.⁷ Unlike traditional nanoscale drug delivery strategies, nanobots offer a whole range of possibilities that include targeting design perspective and the ability to sense, control, and carry out massive tasks in parallel. For instance, remote controlling by manipulation of magnetic and optical properties in an origami-like fabrication of microbots can provide excellent theranostic applications for several brain disorders.⁸ Thus, extending the existing nanodrug delivery strategies to nanorobotics can shape the future of neuromedicine. This review will provide a brief outline on the physiology of BBB that is significant in in vitro model development. The review further proceeds with explaining the existing model system that a nanoroboticist can utilize to design nanobots for targeting BBB with maneuverable properties. Finally, an attempt has been made to summarize the recent advances in the nanorobotics for BBB crossing theranostic application by utilizing computational modeling and the technological gaps that needs to be filled by future researchers.

1.1. Early Attempts To Identify BBB Existence and Establishing Its Relation with Central Nervous System (CNS). Blood vessels are the primary infrastructure involved in the transport and delivery of oxygen and nutrients to all organs. Being the most complex element of human body, the brain demands a more complex microvasculature. Research on this dynamic conduit has been ongoing for two centuries with several significant findings (Figure 1). Though the original theory on the existence of a barrier that prevents movement of molecules was identified by Paul Ehrlich based on dye injection studies, the term "blood-brain barrier" was initially coined by biochemist Lina Stern, after the systematic study of transport of several molecules from blood to brain.⁹ Similar to studies by Stern and Gautier, several investigations were carried out by dye injections to understand the circulations into the human brain.¹⁰ The next important milestone on BBB was the identification of cellular structure that was responsible for its barrier properties. After the advent of electron microscopy, visualization of BBB became feasible and attempts were made to understand the cellular architecture of BBB (Figure 1). By injecting horseradish peroxidase, Reese and Karnovsky¹¹ evidenced the confinement of the same luminal region which was further attributed to the tight junctions between two adjacent endothelial cell (EC) membranes. Further investigations on BBB were focused on identifying the phenotype involved in the barrier. Grafting experiments performed in vitro threw light on the involvement of astrocyte endfeet and several other cell types in the barrier function.¹⁰ Early in vitro studies by Arthur and team¹² revealed the existence of tight junction facilitated by the coculturing of endothelial cells with astrocytes. Simultaneous search on other cell types involved in BBB was also conducted by several researchers in the early 19th century. The contribution of

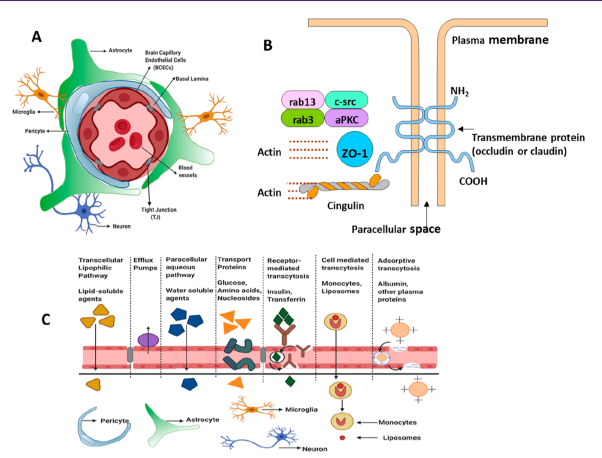


Figure 2. Molecular and transport details of BBB: (A) blood-brain barrier formed by the endothelial cells and contributed by pericytes, astrocytes, microglia, and neurons; (B) structure and function of tight junctions and some of the molecules that contribute to the tight junction; (C) primary transport routes across the BBB. Created with BioRender.com.

barrier properties by pericytes was first identified in the late 19th century by Eberth.¹³ Though there was little research on pericytes in 20th century, emergence of advanced and accurate techniques has influenced the increasing research on these cell types in the past 15 years.

1.2. Complexity in Structural and Functional Anatomy of BBB. On the basis of these initial studies, the current knowledge of the components involved in BBB and their functions has been derived. The BBB is a combination of several cells including specialized EC, along with microvessels, astrocytes, pericytes.¹⁴ All these cells interact continuously and further form the basal membrane of the vascular unit (Figure 2A). The interaction of ECs with other neural and immune cells resulting in an integrated network is known as the neurovascular unit.^{15,16} EC and pericytes and astrocytes contribute to the brain microvasculature and regulate the passage of substances between brain and blood, thus maintaining brain homeostasis and also prevention against pathogens and neurotoxins.¹⁷ ECs of central nervous system exhibit unique characteristics unlike other peripheral ECs that can be attributed to the tight junctions (TJs) and adherence junctions (AJs), thus strictly regulating paracellular transport. The TJs are regulated by the TJ proteins composed of claudins, occludin (ZO-1, ZO-2, ZO-3), and junctional adhesion molecules (JAMs) by sustaining TJ structure as shown in Figure 2B.¹⁸ Keaney and Campbell¹⁹ noted that AJs play a role in maintaining TJs and junctional complex. While TJs control the intracellular transport based on molecular

properties including ionization, lipophilicity, polarity, and other physicochemical properties, the intercellular transport is facilitated by diffusion, endocytosis, and the ratio of influx and efflux transporters (Figure 2B). Similarly, synaptic signaling is regulated by maintaining optimum ionic composition using ion channels and such transporters.^{17,18} Despite the existence of several literature works on the molecules involved in TJ and their role, specific AJ function is yet to be studied in depth.

The research on astrocytes contribution in BBB has increased ever since its role was identified by grafting experiments. Further coculturing with astrocytes induced formation of tight endothelium with higher trans endothelial electrical resistance (TEER) values, thereby reducing the permeability of different molecules.^{17,20} Anatomically, pericytes are part of the abluminal surface (Figure 2A), embedded within the basal membranes.²¹ Though the exact role of pericytes in barrier properties is still unknown, the relatively higher ratio of pericytes to EC in the central nervous system is considered to be responsible for the endothelial barrier properties. Similarly, neurons and microglia further contribute to maintaining the integrity and functions of BBB. The transport of molecules is highly controlled by this unique microvasculature. With the lack of evidence on the exact function and properties of other cell types, there is a wide scope for research on studying the role of all cell types during health and diseased state. With the combined activity of all BBB cells, resistance to paracellular diffusion of ions and other

Table 1. Entities Crossing the BBB

	active transport	passive transport	mixed transport	remark
Lipid-soluble molecules		yes		Influenced by concentration gradient and lipid solubility
				Example: blood gases and anesthetics, heroin ²⁵
Solute carriers (SLC)	yes		yes	Driven by electrochemical (i.e., Na^+ or H^+ pump) or by concentration gradients ²⁶
				Example: drugs like L-DOPA, glucose, amino acids, nucleosides, monocarboxylates, and organic anions and cations
Carrier-mediated efflux (efflux transporters)			yes	Involves energy expenditure by ATP hydrolysis for transport against concentration gradient
				P-glycoprotein (Pgp, ABCB1) and breast cancer related protein (BCRP, ABCG2) are some efflux transporters
				For example, a variety of cytotoxic drugs ²⁷
Receptor mediated transport	yes			Induced by binding of molecules to receptors like insulin receptor, low density lipoprotein (LDL), transferring receptor
				Example: nutrients like insulin, iron, and leptin ^{28,29}

molecules resulting in high TEER can be noted.¹⁸ As a result, all the cell types and the BBB properties become highly pertinent with respect to treatment, diagnosis, and imaging for brain abnormalities.

While TJs maintain homeostasis by restricting passage of toxins and pathogens, the energy and nutritional requirement for the brain is achieved by several specific transport mechanisms, viz., paracellular, transcellular pathways (receptor mediated and carrier mediated), adsorptive transcytosis, and passive diffusion (Figure 2C). Small lipophilic molecules like carbon dioxide are transported across the BBB by passive diffusion. The mechanism of passage of components between the endothelial cells is termed as paracellular transport, whereas the passage of molecules through the endothelial cells is achieved by transcellular transport. In a healthy brain, a proper balance of paracellular and transcellular is decisive for determining the permeability properties.²² The endothelial cell lining of BBB is known to possess specific receptors for plasma protein growth factors and several macromolecules on the luminal and basolateral side of the ECs (Figure 2C). For example, glucose transporter 1 (GLUT-1), neutral amino acid transporter, and cation and anion transporters are known to play key roles in brain metabolism. And this is usually achieved by carrier mediated transport of their substrates.²³ While some transport pathways are simply governed by a concentration gradient, some require energy for the transport of the desired molecules to the brain. In such cases, adenosine triphosphate (ATP)-driver efflux pumps (also known as ATP binding cassette (ABC) transporters) contribute to molecular passage across the BBB to maintain brain homeostasis (Table 1). Neurotoxins are primarily excluded from the brain by this passage. Active pharmaceutical ingredients (APIs) also become the substrate of these transporters and get excreted. Some of the extensively researched efflux proteins are the Pglycoproteins (P-gp), breast cancer resistant proteins, and multidrug resistance associated proteins (Table 1).

Some molecules do not have specific transporter proteins but still manage to reach the brain. Such transport is usually facilitated by receptor mediated transcytosis. This is achieved in three steps, viz., endocytosis, intracellular vesicular trafficking, and exocytosis. Molecules that need to be transported get attached to their respective receptors on the endothelial cell lining, thereby initiating endocytosis. This molecule and receptor complex are invaginated leading to the formation of intracellular transport vesicles. While the receptor gets recycled, the molecules get sorted and exocytosis is carried out by release of the molecule at the basolateral side.²³ Insulin receptors, low density lipoprotein receptors, and leptin and lactoferrin receptors are some of the receptors that carry out receptor mediated transcytosis. A more detailed review on the mechanisms of transport across the BBB has been reviewed elsewhere.²³ Another mechanism by which molecules cross BBB is by adsorptive mediated transcytosis which is a nonspecific pathway unlike a receptor mediated one. Though this mode of transport exhibits low affinity, it shows higher binding ability, thus having similar transcytosis efficiency.²⁴

A number of neurological conditions including multiple sclerosis (MS), stroke, epilepsy, vascular dementia, Alzheimer's disease (AD), Parkinson's disease (PD), brain infections, and various neurological tumors can be attributed to the dysfunction of BBB. In all these disorders, BBB dysfunction is the primary element of pathology influenced by a series of physiological properties. For instance, basal membrane degradation, altered expression of components in efflux pumps, and leaky BBB occur in series, leading to brain disorders.³⁰ Similarly, in the case of traumatic brain injuries, such events occur immediately or in a delayed manner and result in inflammation and activation of coagulation cascades. Recent evidence also suggests that poor expression of TJ proteins is the underlying reason for several psychiatric disorders.³¹ Similarly in the case of MS pathology leaky barrier was attributed to being the underlying reason for lesion formation and disease progress.¹⁶ It is eminent to mention that in many diseases, it is unclear whether disease is caused due to BBB dysfunction or disruption in BBB is the result of disease. Thus, a thorough understanding of the BBB properties will facilitate better model designs and targeted thernostic development.

2. *IN VITRO* MODELING OF THE NEUROVASCULAR UNIT: A CRITICAL APPRAISAL OF ADVANCES IN THE FIELD

Successful replication of the neurovascular unit provides an invaluable tool to aid in dissecting out the pathological factors, mechanisms of action, and the onset of CNS disorders. Models are crucial to predict the uptake of drug candidates prior to costly and laborious in vivo studies. The human body is composed of both cellular and noncellular material organized in a highly specialized manner. It is difficult to mimic all aspects of human biology with one in vitro model system. Nevertheless, in vitro models continue to hold significance in

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brain research, drug development, and testing owing to their flexibility in designs.

2.1. Primary Cells, Cell Lines, and Human-Induced Pluripotent Stem Cells Model. The first and foremost challenge in developing a BBB-on-chip is the choice of cell lines or types used in the model development, specifically brain microvascular ECs. It will determine the translational relevance of the model since the model exhibiting highest similarity to in vivo BBB structure and functions is considered to be a more reliable model. Primary cell lines are regarded as the ideal candidate, as they provide the closest approximation to their in situ counterpart.³² Such primary and low passage EC has the ability to mimic or reproduce maximum biochemical functionalities and possess morphological similarities to their in vivo counterpart.³³

Initially, several BBB models were established by culturing primary human brain ECs, by isolating the tissues during brain specimen resection in tumor patients or during autopsy. However, concerns regarding the availability, poor yield, contamination by other cell types, and reproducibility of the cell lines limit their ubiquitous use in BBB models.³³ On removal from the host environment, human brain ECs dedifferentiate rapidly and lose their primary BBB properties, leading to drift in the phenotypic traits resulting in leaky paracellular function.³⁴ To circumvent this problem, cell immortalization was attempted by few researchers. Human cerebral microvascular endothelial cell line (hCMEC/D3) is one of the most commonly used human immortalized cell line in the BBB model development other than immortalized cell lines from animals.³³ However, few reported poor tight junction protein expression resulting in low TEER values, thus making them unworkable for BBB model development.³ Alternatively, primary brain ECs from mouse, bovine, rat, and porcine like mouse b.End.3 cell line, rat GP8, RBE4 cell line, etc. have been used for developing the in vitro BBB model as they pose better advantages than primary cell culture in terms of the ability to form better functional BBB, low cost maintenance, and faster growth.³⁶ As the brain microvasculature occupies only 0.1% (v/v) of the whole brain, a large number of rodents are needed to generate the required brain ECs. Nevertheless, similar to in vivo models, species difference results in poor translational relevance.³⁷ For example, it was noted that there was a dramatic decrease in the activity of γ -glutamyl transpeptidase and alkaline phosphatase by the RBE4 cell line in comparison with isolated brain capillaries.³

Identification of potentials of pluripotent stem cell (PSC) came as a breakthrough for development of model systems of human and has cleared a path for novel avenues in the field of neurotherapeutics. These preimplanted embryo-derived renewable cell lines have inherited developmental capacity to generate all kinds of cell types.³⁹ They are known for their ability to renew by itself and differentiate into phenotype by pluripotent. Researchers have ever since attempted to differentiate pure population of ECs from PSC that are capable of possessing BBB properties.⁴⁰ Human induced PSCs possess several key characteristics of brain microvascular ECs including good tight junction properties and appropriate expression of efflux transporters, thereby having an effective measurable TEER value.⁴⁰ Hence, a large accumulation of literature in recent years on BBB models involving these cell types can be seen. Many researchers have also identified enhanced barrier function by coculturing induced PSC based brain ECs with primary astrocytes, pericytes, and neural cells.^{41–43} Further focus on using induced PSC derived cell types of astrocytes and pericytes was also attempted recently to note an improvement in the TEER values,^{44,45} thus making a personalized approach on modeling BBB, with the hopes of gaining new insights on genetically influenced neurological disorders. Use of models with induced PSC offers potential for investigation of BBB breakdown at the earliest stages of dysfunction, which can be difficult from post-mortem tissue use.⁴⁶ However, reports have identified the lack of complete translational relevance of iPSC based brain ECs to their in vivo counterparts like showing epithelial markers along with endothelial cells.⁴⁷

2.2. Components of Microengineered Perfusion Based Preliminary BBB Models (Structural Basis). In vitro modeling of BBB has seen many radical changes and novel technologies as an amelioration of the existing traditional techniques. Several in vitro BBB models have been developed including monolayer, coculture, dynamic models, and microfluidic models. It is to be noted that no in vitro model can exactly replicate the in vivo conditions. However, they bridge the gap between pathology and drug discovery. By understanding the limitations of the different models, a better model can be designed and applicability of the data obtained can be determined by better assessment of the results. BBB models involving cell cultures are the most versatile model for basic research on permeability studies.⁴⁸ Cell culture models, based on either primary cells or immortalized brain endothelial cell lines, have been developed in order to facilitate the in vitro studies of drug transport to the brain and studies of endothelial cell biology and pathophysiology. One primary challenge in these models of BBB is that brain ECs undergo rapid dedifferentiation when removed from their native environment, resulting in a generic phenotype.⁴⁹

These in vitro models can be classified as static and dynamic models based on their ability to create shear stress mimicking physical traits of BBB. Static cultures are employed in studying transport kinetics, signaling pathways, and high throughput screenings.⁵⁰ Most of the static BBB models employ cell cultures in Transwell systems. On the basis of the number of cell types used, static model can further be classified as monolayer model and coculture system. In the former, the brain ECs are cultured on top of a permeable membrane, thus enabling compartmentalization of blood side and brain side. This microporous membrane actively prevents migration of cells to the other side while selectively permeating only the small molecules or cell expressed protein components. To maintain the selective permeability and nonspecific diffusion, the BBB associated in vivo capillaries are generally 1-3 cell layer thick, and in vitro experiments also demonstrate that cell viability is preserved in 40 μ m or 1–3 cell thick layer.⁵¹ Therefore, Transwell membranes which are $10-30 \ \mu m$ thick exhibit a suitable in vitro culture system for the BBB experiments. A more detailed review on the types of porous membranes adapted for this application has been discussed elsewhere.⁵² Due to the polar nature of the brain ECs, apical blood compartment and basolateral brain side are achieved. It is to be noted that the monolayer BBB model has only one cell type, the brain EC, and has no communication with any other cell types. However, there is more room for maneuvering in Transwell setups, thus making coculturing with other cell types feasible.53 Due to such flexibility in its design, Transwell systems still continue to be widely used as one of the primary

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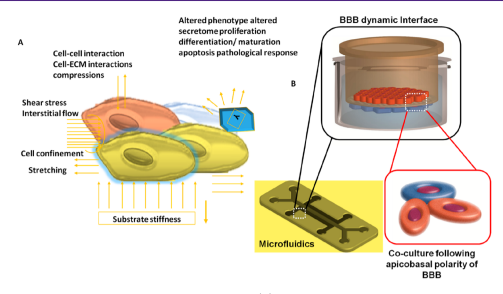


Figure 3. Physical traits and dynamic environment in and around BBB. (A) Principle physical forces ensuring the barrier function of BBB; (B) cutting edge in vitro BBB models following perfusions system and BBB specific cytometry to recapitulate the advanced BBB models.

BBB models, owing to their user friendliness and ease to set up. However, there are several limitations inherent to these models including 2D structure and poor or no exposure of EC to shear stress.

3D cell culture is of representation that is more accurate. In contrast to static models, dynamic models were created with the sole purpose of replicating the physiological surrounding of BBB and addressing the limitations of 2D cultures in terms of in vivo tissue specific functions. A range of approaches have been used to create shear stress in BBB models including hollow fiber constructs (spheroid model) and microfluidics systems.⁵⁴ Such dynamic in vitro BBB (DIV-BBB) holds the ability to self-assemble to resemble the anatomy of brain microvasculature with high TEER values.⁵⁵ DIV-BBBs are one of the initial 3D systems developed to replicate the complexity of the BBB with ECs cultured on the luminal side of the capillaries or hollow fibers, while glial cells are seeded on the outer surface.^{56,57} A comparative study between cell barriers cultured in a static flow (Transwell) and dynamic flow (DIV-BBB) environments showed that fluid flow conditions greatly influence how a cell barrier develops. TEERs across the DIV-BBB culture increased to 10-fold higher than those recorded in the Transwell and are much closer to values observed in vivo.⁵

2.3. Organoid and Spheroid in Vitro Models. Research with a more realistic 3D model of BBB can better mimic the BBB properties in vivo. For this purposed 3D cultures can be classified based on the complexity of the structure as spheroid, multicellular spheroid, and organoids.⁵⁸ On the basis of the culture conditions and methods, different models can be achieved. Though many organs have been modeled using these techniques, research on BBB spheroids and organoids is very limited owing to their complexity. Organoids are in vitro derived 3D cell aggregates derived from primary tissue or stem cells that are self-renewed and self-organized and exhibit organ functionality. BBB organoids are obtained following coculture of endothelial cells, pericytes, and astrocytes under low adhesion conditions. Such a self-assembling model provides a certain degree of developmental freedom resulting in a model of BBB structure with similar complexity.^{59,60} The noteworthy feature of organoids model is that each cell type in the organoid is in direct contact with each other which is crucial

for BBB integrity. As a result, organoids reproduce many features of the BBB, including the expression of tight junctions, molecular transporters, and drug efflux pumps and hence can be used to model drug transport across BBB. They are one of the relevant models of in vivo conditions and proved to be a stable system for extended cultivation. They are a cost-effective and versatile model for therapeutic discovery for the treatment of various neuropathologies. Research on cerebral organoids has seen significant advances in recent years with more specific protocols for developing vascularized organoids.^{61,62} One of the critical drawbacks of organoids is the absence of essential types of cells, including glia, microglia, oligodendrocytes, vasculature, etc. This may hinder neurons maturation, thus limiting its utilities for specific disease models.

To overcome the limitations posed by organoids, 3D spheroid model of the BBB comprising all major cell types including neurons, microglia, and oligodendrocytes were developed to recapitulate more closely normal human brain tissue. Spheroids show expression of tight junctions, adherent junctions, adherent junction-associated proteins, and specific markers. Use of induced pluripotent cells to derive the cells could help narrow the gap in achieving an ideal in vitro BBB model.⁶³ For example, high cell viability was found to be maintained up to 21 days in the spheroid model containing six cell types, which is useful in evaluating long-term effects of drug toxicity. Expression of P-gp and GLUT-1 proteins was identified. Junctional protein distribution was altered under hypoxic conditions.^{64,65} Thus, the spheroid model may have potential applications in drug discovery, disease modeling neurotoxicity and cytotoxicity testing. However, structural consideration in terms of extracellular matrix and cell specific function needs to be addressed to make this model applicable for pathology studies.

2.4. Microfluidics and Application of 3D Printing. The behavior of vascular cells is greatly influenced by mechanotransduction creating the ability to convert biophysical stimuli to biochemical signals in the cellular microenvironment.⁶⁶ Due blood flow, brain ECs are subjected to shear stress that in turn realigns and elongates according to flow by redistribution of junctional proteins (Figure 3). Additionally, the mechanism involving vascular endothelial (VE)-cadherin, platelet endo-

Table 2. Current Model of in Vitro BBB with Merits and Demerits

type of model	features	advantages	disadvantages	
Static 2D				
• EC monolayer on Petri dish	Endothelial cells grown directly on Petri dishes filled with medium	Cheap and facile standardization	Quick dedifferentiation of cells	
		High throughput screening	No shear	
			Limited barrier properties	
			Can be applicable only for monolayer	
			Lack of cell-cell interaction	
Static Transwell 2D				
• Monolayer	Endothelial cells gown on porous membrane	Cheap and easy standardization	Lack of cell-cell interaction	
			No shear	
• Coculture	Endothelial cells and astrocytes and/or pericytes	EC-perivascular interaction	Lack of blood flow and shear stress	
	grown on either side of porous membrane to form apical and basal side		Limited barrier properties	
	apical and basal side		Limited cell-cell interaction	
3D				
• DIV-BBB	Initial models of 3D culture	Improved physical attributes	Requires high cell numbers	
		Low cost fabrication	Invisible cell-cell interaction	
		Enable shear stress with capillary- like structures	Limited leukocyte transmigration	
		Possible hemodynamic studies	Not ideal for high throughput screening and kinetic studies	
• Organoids	Self-assembling under low adhesion of EC,	Can be made patient specific	Lack vasculature	
	astrocytes, pericytes	In vivo like complexity and architecture	Less amenable for high throughput screening	
			Assay complication	
• Spheroids	Involves microglia in addition to ECs, astrocytes,	Ease of use	Simplified architecture for	
	pericytes	High reproducibility and scalability	pathological studies	
 Microfluidics BBB 	Traditional microfluidics patterns by soft lithography	Flexibility in design	Moderate TEER	
	with channels for coculture of EC and astrocytes/ pericytes	Less cell number requirement	Limited scalability	
		Control of microenvironment	Complex fabrication	
		Considerable shear stress		
		In vivo like structure		
• 3D printing	Patterning of the channels by 3D printing similar to	In vivo like architecture	Lack of high through put screening	
	in vivo microenvironment	Immediate permeability studies	Not ideal for kinetic studies	
		Flexibility in design		
		Built-in multicellular network		

thelial cell adhesion molecule (PECAM-1), vascular endothelial growth factor (VEGF), etc. is considered to be responsible for this effect by the cells.⁶⁷ With the aim of recreating this intricate biomechanical interaction, application of microfluidic devices was adopted giving rise to a new generation of BBB models "brain-on-chip". Due to its several advantages, microfluidics has emerged as the innovative approach for conducting research related to the brain, including modeling of neurodegenerative diseases and high throughput drug screening. A typical microfluidic pattern is derived by soft lithography of elastomeric material, and the cells are cultured on the channels under dynamic flow to induce cell proliferation. These so-called "organ-on-chips" contain microfluidic devices, in which cells are cultures in a continuously perfused, micrometer sized compartments to mimic the physiological conditions in the tissues. The different aspects of microfluidics and their application in brain research have been discussed in several reviews.^{54,67,68} To date, several designs of microfluidic BBB exist, including sandwich, parallel, 3D tubular design, and vasculogenesis pattern.⁶⁹ The former two are a more preliminary and well explored model, while the latter are the advanced design of microfluidic models. As a recent advancement in the microfluidic BBB on chip development, 3D printing to replicate the neurovascular microenvironment by collagen was attempted to conduct studies on drug screening

and inflammation.⁷⁰ By control of the cell patterning, different cell types can be cultured in distinct patterns similar to their environment in vivo. 3D printing of microfluidics network highlights the most recent advances in BBB-on-chip devices, where elements are added to fabricate objects based on 3D model data. The intended product is digitally rendered in 3D with computer aided design (CAD) software. Raw materials such as thermoplastic polymers, natural polymers, and biocompatible synthetic polymers are processed into filaments and granules. Unprinted materials will also be harvested and recycled for continued use in the printing process. The leading 3D setup processes for microfluidic systems are 3D printed transfer molding, fused deposit in modeling, stereolithography, direct ink jet printing, 71 and selective laser sintering. Polyjet,⁷³ digital light processing,⁷⁴ liquid deposition modeling,75 and fiber encapsulation additive manufacturing are the new developed techniques.⁷⁶ Despite the several advantages of 3D printing in BBB microfluidics, the technology is yet to gain momentum due to several bottlenecks. For instance, the complexity of the tissue to be reproduced increases exponentially the complexity of the technical challenges that need to be overcome, thus hindering the widespread adoption. However, complete integration between multiple fields and examination of validation steps will strengthen their future applications. The key attributes with merits and demerits of pubs.acs.org/chemneuro

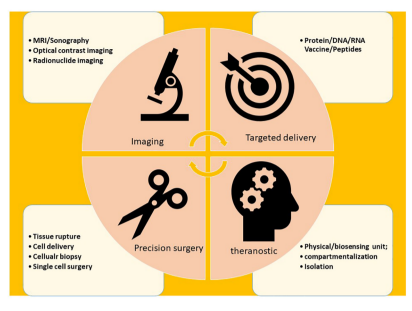


Figure 4. Emerging scope of nanorobotics in neurology. The four quadrants of the scheme demonstrate the trend of nanorobots application in imaging, nanomedicine, nanotheranostic, and envisioned nanosurgery in single targeted cells, a visionary cutting-edge technology for precision medicine with versatile task achievement.

each model are listed in Table 2 to facilitate wise choice of models for future theranostic development. Despite the several advances in different in vitro models, there continues to be prospects for newer and more advanced models, aiming at tighter junctions, close cell–cell interaction, and better shear induced BBB morphology, ultimately aiming at close mimicking of BBB properties. With very limited therapeutic options for neurological disorders, in vitro modeling of BBB continues to appeal to a majority of neuroscientists.

3. ADVANCES IN NANOROBOTICS-ENABLED TARGETED DRUG DELIVERY ACROSS BBB

The primary drive in the field of neuromedicine is to develop smaller, efficacious, and cost-effective systems that will easily cross the BBB and reach the brain parenchyma. Such a quest is based on the cellular and subcellular genesis of neurological disorders. Of the different approaches available to circumvent the BBB defense mechanisms, application of nanorobotics has gained more popularity due to their multifunctional approach. The programmed nanoparticle to theranostic delivery across BBB can be addressed as "nanorobots or nanobots" since they are capable of carrying out specific tasks with controlled maneuvering and targeting." Nanorobots are biocompatible entities of size 1-1000 nm and are synthesized using several biomaterials including lipids, polymers, metals, and crystals. Scaling down the tiny robots from microscale to nanoscale reveals several unique scale-dependent properties such as low power independent actuation and dynamic switchability between 2D and 3D swimming.⁷

The recent focus on controlling and maneuvering the nanorobots in in vivo microenvironment has attained major thrust in the field of theranostic application. In this context, the control of nanorobots requires device modeling (e.g., biohybrid, self-(dis)assembly driven), fields (e.g., untethered magnetic field, optical, electric field), and feedback (e.g., open or closed loop). In such a case, manipulation of nanobots by external magnetic field in the clinical environment using magnetic resonance imaging (MRI) will be feasible and can be precisely relocated for delivery in the brain. The magnetic torque applied by uniform magnetic field lines to nanorobots aligns and redirects the nanorobots motion, providing a high degree of local control on device and facilitating long lasting operation.⁷⁹ Further control and maneuvering of bacteria driven nanorobots could be harnessed in the future, riding upon advances in synthetic biology to produce programmable and functional magnetic components via genetic manipulation responsive to local environment.^{80,81} Similarly, optical control is another practical possibility, with limited penetration to body tissue, and is still in a state of infancy for nanorobots control. The closed loop control feedback, often called feedback control systems, is the method of choice with the ability to self-correct in an autonomous in vivo microenvironment.⁸²

Nanomaterials can be formed in different structures with specific size and other physical properties like optical, photodynamic, magnetic, etc. Owing to their flexible properties, they can be used to target specific sites. The nanorobots are tamed to selectively sneak through the permeable membrane, which regulates the passage of a multitude of large and small molecules into the microenvironment of the neuron, without perturbing the homeostasis of CNS. As shown in Figure 4, these stealth nanobots are selectively programmed to disguise those traffic posts via molecular mimicry to assist a multitude of tasks such as imaging, delivery, sending, and precision surgery.⁸³ Bioengineers and roboticists are developing ways to safely bypass the BBB to deliver the therapeutics to the brain without any long-term effects. In this regard, nanotechnology is providing the permeable platform enabling delivery of drug across the BBB. For example, nanodiamonds, a less toxic substitute to other carbon nanoparticles, are currently applied in biomedical imaging, drug delivery, and other areas of medicines.⁸⁴ The excellent biocompatibility, functional versatility, and unique surface electrostatic properties of nanodiamonds are attributed to the drug delivery.

Regardless of the high restriction to the transport of molecules across the BBB, the CNS continues to demand

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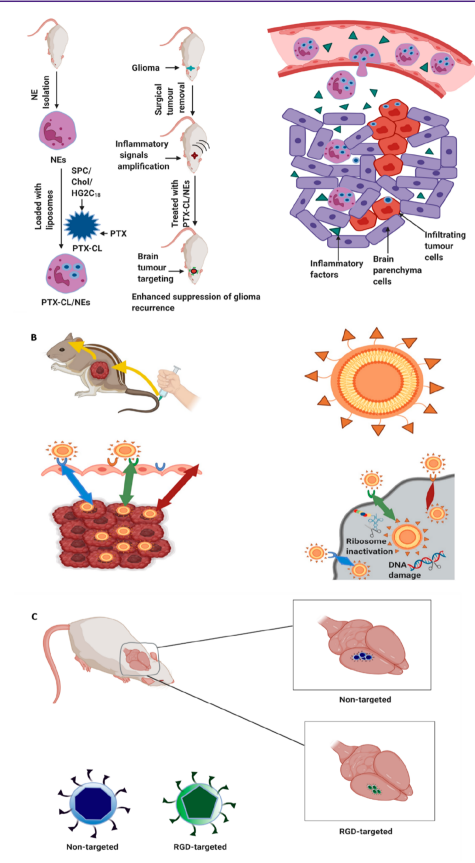


Figure 5. Multifunctional design criteria of nanobots to cross the BBB: (A) varying ApoE ligand density to tame the barrier transport of nanobots for therapeutic delivery scheme involving multireceptor mediated therapy; (B) soft-nanobots decorated with paclitaxel-loaded cationic liposome targeting to suppress postoperative glioma recurrence crossing BBB; (C) schematic illustration of nanobots with integrin based sorting and detection of brain glioblastoma comprising cell penetrating RGD peptides for surface enhanced Raman spectroscopy. Figure is created with BioRender.com.

nutrients and energy to be regularly supplied to it. For this purpose, several mechanisms exist that facilitate the transport of small molecules that keep the CNS functions intact. To effectively deliver drug molecules, one should tap into the potential of these transport mechanisms, thereby leading to an effective therapeutic opportunity without having to succumb to brute force techniques that involve the mechanical or physical disruption of barrier.⁸⁵ Nanobots are designed to be able to cross the blood-brain barrier through several pathways to enable communication and transport of nutrients across the blood-brain barrier in endothelial cells. The prospect of delivering drug by passive diffusion is very limited due to the highly selective nature of BBB. This is arbitrated by the TJs and adherence junctions in addition to the phospholipid part of ECs. The only possibility for drug delivery by this mechanism is to develop a small lipophilic molecule, which in turn is highly dependent on the surface area, charge, molecular weight, and volume.⁸⁶ With higher lipophilicity, lower hydrogen bonding further reduces the diffusion.⁸⁷ Thus, net migration of components is highly dependent on the concentration gradient, thereby becoming a poor choice of transport pathway for drug molecules, especially through nanoparticulate transport.¹

By specific design of the nanoparticle to mimic the components that have the ability to undergo paracellular transport, targeted drug delivery can be achieved. Also, utilizing the potentials of components naturally entering BBB by this pathway can also be explored for targeted theranostics. For instance, neutrophils have the natural ability to cross the BBB and infiltrate the tumor mass in a brain cancer patient. These immune cells can be used as "Trojan horses" to deliver the drug payload or imaging agents to the affected sites. Xue and team⁸⁸ explored the ability of neutrophils to deliver paclitaxel loaded cationic liposomes (PTX-CL) into the inflamed brain tissue after surgical resection. It was noted that such a designed PTC-CL/NE delivery system significantly reduced the recurrence of the brain tumor (Figure 5A).

It is to be noted that many of the targeted drug delivery using nanoparticles focuses on crossing the BBB through transcytosis, owing to the ability to develop nanocarriers specific to receptors.^{89,90} Polymer and magnetic nanoparticles facilitate drug delivery across the BBB through receptor mediated transcytosis.^{3,91} Nanocarriers can also be designed to target receptor mediated pathway by preparing the nanocarriers-like extracellular vesicles secreted by cells so that they can carry the drug load across the BBB through endocytic pathways.⁹² By utilization of the potential of ligands that can facilitate the crossing of BBB via carrier mediated or receptor, mediated pathways can effectively release the desired compounds, thereby facilitating drug delivery or enhanced site-specific imaging feasibility. For example, the BBB crossing ability of apolipoprotein E peptide (ApoE) and SAP loaded chimeric polymerosome was investigated.⁹³ It was noticed that ApoE promoted multireceptor mediated pathway including LDLR, LRP1, and LRP2 (Figure 5B). In order to develop efficient nanobots, one should know the different mechanisms by which molecules get transported across the BBB. By tailormaking nanomachines, a different transport mechanism can be achieved with a wholesome theranostic application.

Brain cancer or glioblastoma is highly invasive and one of the most devastating deadly neoplasms. The prognosis remains dismal, and the median survival rarely exceeds 16 months. The present clinical treatment approaches include surgery, chemotherapy, and radiotherapy. However, it is widely known that complete surgical ablation is impossible and the possibility of recurrence is high.⁹⁴ Nanotechnology poses as a remarkable alternative to such a conventional, invasive diagnosis and treatment in the field of brain tumor. For example several fluorescence, photoacoustic, and Raman imaging nanoprobes (Figure 5C) have been highly efficient for intraoperative imaging of tumor cells.^{95,96} A wide spectrum of potential drugs have been investigated to treat several neurological disorders, but their therapeutic success is still limited due to a range of challenges. The past decade has witnessed an unprecedented expansion in engineering of various kinds of theranostic nanobots for cancer imaging and therapy (Table 3). The development of theranostic nanobots that are tumors-specific, safer, simpler, and yet still powerful will continue to be the focus in the near future, holding a greater potential to be translated into the clinic.

Nanomaterials as a possible therapeutic system for targeting neurological disorders has gained more attention due to their versatile properties. Unlike conventional drugs they hold the potential to overcome the BBB defenses, thereby becoming the nucleus of several neurological research. Despite the rapid advancements and the abundant application of nanomedinces, their translation to clinical process has been very slow. To date, very limited nanomedicines have been approved for clinical use by FDA and many nanoparticles are continuing to be the subject of several clinical trials in recent years.¹¹ It is to be noted that the vast majority of the approved nanomedicines and those in development are dedicated toward cancer therapeutics.¹¹² However, the field of nanomedicine is continuing to be the forefront of several research studies, owing to their potential to surpass several bottlenecks, particularly in the field of brain theranostics. Considering the limited number of nanomedicines for neurological disorder to date, an augmented interest toward nanomedicines for overcoming BBB defenses can be expected in the near future. It is to be acknowledged that the development and clinical translation of nanomedicine for neurological dysfunctions are accompanied by their own hitches, owing to the limitations posed by the size and also the recent necessities to develop safer nanomaterials.¹¹³ This can be attributed to the poor predictability of these nanomaterials' behavior and state in the complex BBB atmosphere. For instance, disease pathology in each patient may vary, thus affecting the effectiveness of the nanoparticle. Similarly, the drug behavior with humans could vary from their behavior in animal models with the same pathological condition resulting in failure of clinical translation due to such discrepancies. Recent advances in bioprospecting of natural resources for development safer and natural nanotherapeutics could hold the future of BBB targeting nanomedicine.^{114,115} Further investigations on the nanomedicine properties like their pharmacokinetics, pharmacodynamics, and bioavailability need to be thoroughly studied in addition to their toxicity, side effects, and biocompatibility prior to clinical use. For example, the tamed nanorobots after extravasation from the blood vessels will largely be trapped into the cells in closest distribution of blood vessels (Figure 6). Furthermore, even if the nanoparticle meets the standards of clinical trials, at a manufacturing point of view, the scale-up of nanoparticle synthesis is difficult, making them a poor choice for pharmaceutical companies to venture into bulk production. Despite the challenges in nanoparticle development, the promise that nanomedicines hold in the field of brain

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nanocarriers	drugs delivered	disease/disorder targeted	effects	nanotechnology assistance and control mechanism	ref
Solid lipid nanoparticles (SLNs)	Quercetin	Alzheimer's disease	 Ability to reverse degenerative effects of the free radicals induced by exposure to aluminum chloride 	 Prolonged half-life and drug release on demand 	67
			• Improved antioxidant properties of the drug due to encapsulation	• Size dependence control	
Chitosan nanoparticles	Dopamine	Parkinson's disease	• Resulted in a pulsating release of dopamine into the stratum	 Prolonged bioavailability and targeted delivery 	98
				 Can be radiolabeled with radionucleotides for imaging 	
				• Size control dependent control	
Polylactic acid-co-glycolic acid (PGLA)	Levodopa	Parkinson's disease	 Achieved high target tissue concentration with less doparnine levels in blood indicating targeted delivery 	Higher selectivity/sensitivity in targeted delivery	66
	MEM-PEG-PGLA	Alzheimer's disease	\bullet Reduction in eta -amyloid plaques and disruption of the aggregates		100
Redox active nitroxide containing nanoparticles (RNPs)	General neuroprotection	Traumatic brain injury	 Possess high reactive oxygen species (ROS) scavenging capability Reduced brain contusion volume 	• Higher selectivity toward lesions	101
			 Increased neuroprotective microglia production 		
Nanoliposomes	Doxorubicin	Tumor	• Tumor suppressive effects on glioblastoma cells were noted.	 Photothermal therapy with high selectivity and sensitivity 	102
			• The cardiotoxicity exhibited by the drug was greatly reduced due to encapsulation in liposome.	• Selectivity in fluorescence imaging of tumor margin with higher precision	
				• Chemotherapy with higher sensitivity on deep cancer with laser irradiation	
Biomimetic glioma membrane protein liposome		Glioma	 Crossed BBB and showed ability to reach glioma at early stages of development 	 Control possibility by optical field 	103
Cu-labeled micelles and liposomes	Indocyanin green (ICG)	Glioblastoma	• Glioma exosomes with ICG showed higher accumulation at the site of tumor.	 Higher selectivity with unlimited penetration in positron emission tomography (PET) imaging 	104
				 Control by electrical field 	
Multifunctional lipid nanocapsules	Paclitaxel (PTX)	Glioblastoma	 Ability to act as immunostimulatory, thereby providing long-term immunity 	 Chemo/immunotherapy with higher selectivity and sensitivity 	105
			 Nonmethylated oligonucleotides like CpG mediated immunotherapy achieved with high therapeutic index and reduced long-term relapses 	 Receptor ligand based maneuvering 	
Apolipoprotein E (ApoE) peptide conjugated chimeric polymersome (ApoE-CP)	Saporin	Tumor	• Protein toxin delivery	 Assist in cancer chemotherapy and also in target specific radiotherapy 	93
				• Can be combined with siRNA for co-delivery of drug and gene in chemo/gene therapy	94
H-Ferritin nanocarriers	Doxorubicin	Glioma therapy	• Entry into the BBB via transferrin receptor	• Chemotherapy application with prolonged circulation and half-life in drug delivery	106
			 Nanocarrier accumulated in the lysosomal compartment and all the glioma tumor cells were targeted and killed 	Higher specificity and selectivity in surgical imaging	
				 Enables higher BBB crossing ability Maneuvering by ligand receptors 	
Gold nanoparticles	Prevent aggregation of amyloid β peptides	Alzheimer's disease	• Successful inhibition of aggregation of the A β 42 peptide and crossing of BBB	• Targeted drug delivery using nanotherapeutics with higher affinity	107
			• Showed higher bonding affinity to $A\beta$ 42 with rescue from behavioral impairments	 Magnetic and optical field control 	
Integrin targeted surface enhanced resonance Raman spectroscopy (SERRS) gold nanoparticles		Tumour imaging	 Conjugated with RGD peptide for better visualization and diffuse margins of the main tumor 	 High specificity and ultrahigh sensitivity in Raman imaging with limited special resolution 	95

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Enhanced detection sensitivity at the

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 Improved specificity and bioavailability of drug Controlled by magnetic field, optical field 	 Multimodal imaging with high selectivity Commenses inherent drawhacks 		
• Combined with TAT protein to cross BBB and increase higher uptake by the brain parenchyma	 Conjugation of Pittsburgh compound with multiwalled nanotube Multimodal imaging with high selectivity to increase brain uptake and increase biodistribution of drug Attemnt to have a more clinically available modality in imaging 	• Attempt to have a more cumcany avanable moutany in minging	
Neurodrug delivery	Tumor imaging		
Carbon dots, quantum dots, carbon nanotubes			

theranostics is huge and is rightly considered to be the revolutionary therapeutics of this era due to their versatile nature and tremendous potential.

4. MACHINE LEARNING HELPING IN PREDICTION OF CHEMOTHERAPEUTICS CROSSING BBB

The experimental validation of theranostic agents crossing the BBB to investigate the permeability and drug development targeted to the central nervous system is a very long process. Sometimes it may take a decade and at the end has a very low success rate. This is due to the extraordinary complexity of the brain, the degree of side effects, and most importantly the inadequate precision of the BBB model to investigate in vitro. Therefore, it is of paramount importance to develop prescreening tools for large chemical databases with the aim to test different nanobots. Over the past decade, several computational models have been dedicatedly developed to investigate the BBB permeability including machine learning.¹¹⁶ Several approaches are available in this regard, which are reviewed elsewhere.¹¹⁷ Recently computational nanotoxicology and nanomedicine have made great strides for safer theranostic to biomedical materials designs.^{118,119} By application of appropriate algorithms, prediction of BBB permeability with high accuracy can be achieved. A proper amalgamation of the research knowledge, BBB physiology, and permeability models can result in such accurate predictions, thereby making drug development a rapid process. The computational artificial intelligence (AI) in this context could be categorized as application driven screening in neurooncology, CNS infections, and neurodegenerative disorders such as Alzheimer's, Parkinson, and multiple sclerosis. The computation AI could also assist indirectly BBB related neurotechnology paving the engineered nanomaterials' (ENMs) design for BBB theranostic and imaging, AI for targeting and control maneuvering in CNS arena (Figure 7).

There is a big development in quantitative structure activity/ property relationship (QSAR/QSPR), which is based on database/similarity searching perform systematic evaluation and prediction of whether a molecule could cross the BBB or not. The databases store the info about chemotherapeutics with BBB permeability measured in vivo in animal models as logBBB, i.e., the logarithmic ratio of brain to plasma concentration of test or experimental molecule. The modeling algorithm includes fingerprints molecular similarities of a test compound using distance measuring principal coordinates analysis as input.

Several recent studies on application of QSAR models and AI tools have been established for predicting BBB perme-ability.^{120–123} The machine learning algorithms convert high dimensional data into a lower dimensional vector of coordinates for each molecule such as nearest neighbors, support vector machine, deep neural network, and random forests. Identical pipelines are built for the classification or regressions tree development with neural network and similarity matrix computed via principle component analysis (PCA). Outputs of vector of coordinates only then are predicted as molecular weights and log P variables as a measure of solubility because traditionally these are two important factors that explain why certain molecules cross the barrier; however alone they do not seem to be enough for this classification. The advantage of random forests algorithm is that it allows the analysis of individual variables by looking at their impact in the decrease of accuracy based on the number

Table 3. continued

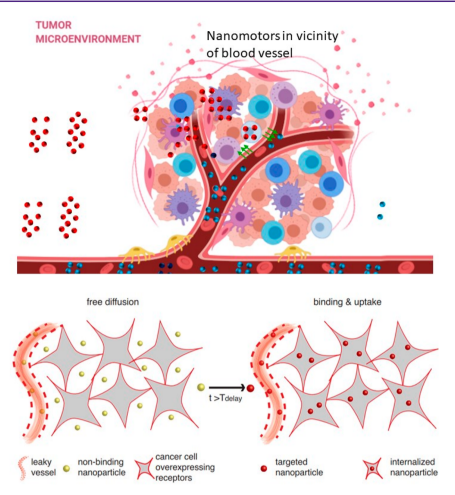


Figure 6. (Upper panel) Enhanced permeation and retention (EPR) effect and nanotheranostic crossing BBB at brain–blood interface showing saturation effect in neighboring microcapillaries (created with BioRender.com). (Lower panel) Delayed-binding strategy. Nanoparticles are shielded from binding for a duration T_{delay} after injection, thereby allowing them to diffuse freely throughout the tumor tissue. After this diffusion period, the shielding is degraded based on pH, enzymatic activity, injected chemicals, or external energy sources to unveil intact targeting ligands. The nanoparticles are then able to bind to the tumor cells. Lower panel is reproduced with permission from Hauert, S., and Bhatia, S. N. Mechanisms of Cooperation in Cancer Nanomedicine: Towards Systems Nanotechnology. *Trends in Biotechnology*. Elsevier Ltd. **2014**, pp 448–455, https://doi.org/10.1016/j.tibtech.2014.06.010.¹³⁰ Copyright 2014 Elsevier.

of bonds in molecular rings as particular descriptors. On the basis of data training, it sets a certain threshold predicating high or low probability of being classified as a crossing molecule. It is expected that the lower weight molecules preferred crossing the BBB. Most importantly random forests choose the number of oxygens with one single bond as the most important variable to distinguish between crossing and then comparison of the overall accuracy. RFs implementation utilizes low P values indicating that data are unlikely with a true null, thus rejecting the null hypothesis for the entire population. This enables high predictive capability for crossing the BBB and the ability to identify the most important molecular descriptors for the particular classification in conclusion of the most potent statistical method for predictions.

Gao and others¹²³ reported significant prediction accuracy gains (from 0.69 AUC to 0.85 AUC) that can be obtained by using both chemical features and clinical phenotypes, compared to using chemical features alone. They have also identified a large number (110) of drugs in a database that can potentially penetrate BBB with their learned model. Subsequent work studies the same problem and improves prediction accuracy using a deep neural network model, in which the neural network is a four-layer multilayer perceptron.¹²⁴ Wang and team¹²⁵ address the data imbalance (i.e., the majority of chemicals in the training data cannot penetrate BBB) through resampling methods, such as SMOTE and SMOTE+. Recently, Alsenan et al.¹²¹ designed and developed a recurrent neural network (RNN) for predicting BBB permeability, which improves prediction accuracy further. The same authors proposed a dimensionality reduction technique Auto-KPCA, which applies kernel principal component analysis (KPCA) as a preprocessing step to enhance the accuracy performance of the subsequent deep learning model.¹²⁶

Finally, it is noteworthy to mention that although the above methods can accurately predict BBB permeability of a given chemical, they do not directly help generate a *de novo* chemical structure with desirable BBB permeability properties. The state of the art for AI-based chemical synthesis¹²⁷ follows the methodology of inverse molecular design¹²⁸ and combines deep reinforcement learning with Monte Carlo tree search (MCTS) to search for a molecular structure with target properties that can be synthesized with known chemical reactions. In contrast, MolGAN¹²⁹ directly generates the graph structure of a new chemical that resembles known drugs, based

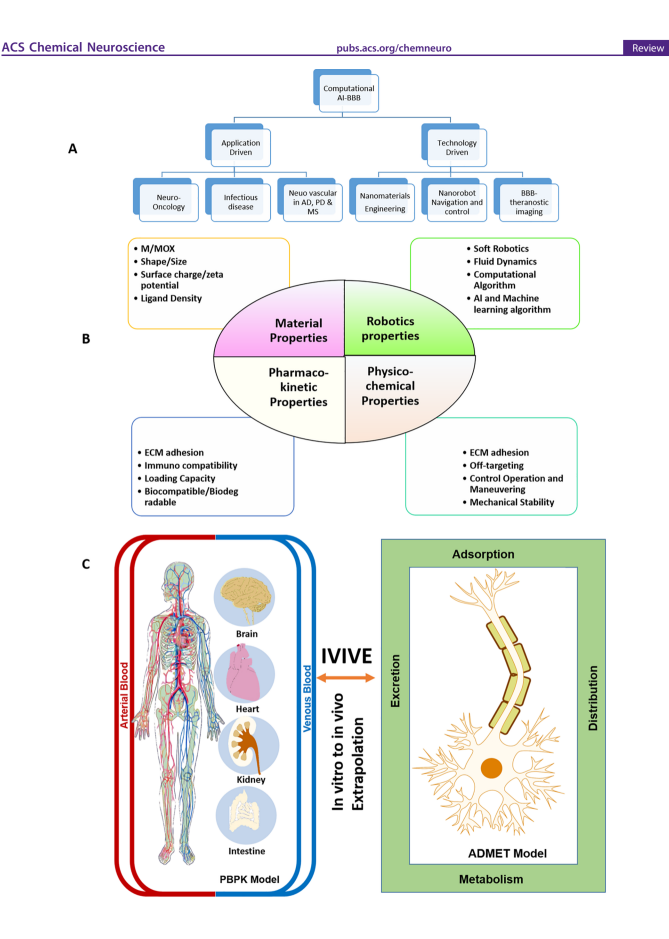


Figure 7. Machine learning and artificial intelligence in advance nanobots design and safety prediction to cross the BBB: (A) scope of computational tools; (B) information driven biophysical design criteria for advance nanobots; (C) physiology based pharmacokinetic (PBPK) models in predicting the safety assessment of chemotherapeutics carrying nanobots for crossing the BBB.

on a generative adversarial network (GAN). None of these methods, however, have been applied to create new chemical structures with BBB permeability, which might be a promising direction for future studies.

5. CONCLUSION AND FUTURE PROSPECT

Navigation across the BBB continues to be one of the primary challenges in neurotheranostics development. While nanodelivery strategies continue to be the best choice in this regard, nanobots can be considered as the pinnacle for surpassing BBB defenses with better control. In future the nanomotors design must accompany application-specific design strategy to follow facile systemic clearance/removal after the therapeutic delivery to the CNS. Therefore, functions required by nanomotors for the therapeutic payload, targeting to BBB, maneuvering control while in systemic circulation and clearance from kidney filtration must be reflected in their physical design intended as per the applications. The smart in vivo performance of nanomotors can be envisioned by selecting self-degradable biopolymers designs that can be broken down into their tiny molecular components to readily clear from brain without causing any immunogenic response or toxicity to brain cells. Therefore, nanorobotics can appreciate plenty of leverage from bioinspired designs strategies involving biomaterials research. In spite of greater development and successful in vitro demonstration of nanorobots with precise control, actuation, and cellular targeting, in vivo applications face tremendous challenges to tame the nanorobots for the theranostic imaging and chemotherapeutic delivery across the BBB. Considering reduction, refinements, and replacement (three Rs) in research, the iterative designs of nanorobots can be improved and tested in the realistic phantoms and ex vivo tissues following scientific ethics and the regulatory permission aspects for rapid bench-to-bedside translation.

Visualizing the dynamics of NPs in the laboratory is often achieved by fluorescent images of adherent cell monolayer culture. Computer simulations have helped model and imagine these complex structures, but stochastic simulations lack physical grounding and are hard to understand. Crowdsourcing test beds can be a key to simplifying stochastic simulations and the physical world predicting homogeneous or inhomogeneous distribution of NPs over cell monolayer depending on tightness of membrane-corona binding. The machine intelligence has to come up with the superior algorithms to tame and control these nanomotors to cross the BBB and reach deeper into the brain tumor. Better algorithms are needed to simulate the binding kinetics, transport, and internalization of nanoparticles on a representative cell monolayer, and stochastic and deterministic reaction-diffusion models can be implemented. Rather than modeling a specific cell line in focus, the experimentalist must collaborate with code developers to focus on a complex scenario whose solution will generalize to a wide variety of cell-NPs interactions addressing in a wide variety of biological barriers environments. Such an advanced algorithm will predict delayed binding to cells close to microcapillaries until the first batch of internalized particles are actively pushed deeper into tumors via active magnetic nanorobots swarms controlled by untethered magnetic coils.¹³⁰

In conclusion, the actual envisioned application of nanorobots to payload delivery across the BBB needs to address the common challenges (synthesis, mobility, tracking, toxicity, biodegradability, etc.) in addition to control and maneuvering schemes. The recent advances in computation design and discovery of nanomaterials using machine learning tools can be utilized to unambiguously address the failsafe synthesis of nanorobots with increasing performance.¹³¹ Similarly nano-QSAR approaches to predict the fate of these nanobots in an in vivo environment could result in effective bench side applications.¹³² However, mobility and tracking in the in vivo environment are a matter of continuous improvement with advancing technologies in design and development of nanorobots to cross the biological barriers. Hence, an integration of in vitro and in silico techniques could result in the development of effective theranostic nanobots for neurological disorders in the future.

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Author Contributions

Conceptualization was done by A.V.S., V.C., S.B., S.P.D., and Y.Y. Data curation was done by A.V.S., V.C., and P.J. Original draft preparation was done by A.V.S., S.B., V.C., and P.J. Review and editing of the manuscript were done by A.V.S., V.C., P.J., A.L., P.L., B.G.-C., J.A., and A.A.A. Graphic design and visualization were done by P.J., D.E.M., and V.C. Supervision was by A.V.S., P.L., and S.P.D. Project administration was done by A.V.S. and S.P.D. Funding acquisition was done by A.V.S., S.B., S.P.D., and A.L. All authors have read and agreed to the published version of the manuscript.

Notes

The authors declare no competing financial interest.

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