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Nanoparticle Brain Delivery: A Guide to Verification Methods

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Nanoparticle brain delivery: a guide to verification methods

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Many reports conclude nanoparticle (NP) brain entry based on bulk brain analysis. Bulk brain includes blood, cerebrospinal fluid and blood vessels within the brain contributing to the blood–brain and blood–cerebrospinal fluid barriers. Considering the brain as neurons, glia and their extracellular space (brain parenchyma), most studies did not show brain parenchymal NP entry. Blood–brain and blood–cerebrospinal fluid barriers anatomy and function are reviewed. Methods demonstrating brain parenchymal NP entry are presented. Results demonstrating bulk brain versus brain parenchymal entry are classified. Studies are reviewed, critiqued and classified to illustrate results demonstrating bulk brain versus parenchymal entry. Brain, blood and peripheral organ NP timecourses are compared and related to brain parenchymal entry evidence suggesting brain NP timecourse informs about brain parenchymal entry.

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Keywords: blood–brain barrier • blood–cerebrospinal fluid barrier • brain parenchyma • capillary depletion • convection-enhanced delivery • in situ brain perfusion • microdialysis • mouse • nanoparticle • rat

The issue addressed in this guide

When considering nanoparticle (NP) distribution, a question that is often raised is whether they enter the brain. With the goal to use NPs to deliver drugs to the brain, many researchers are hopeful that their NPs do enter the brain. Others are concerned about unintended NP brain entry and potential adverse effects. Many have oversimplified the concept of brain entry. They have not considered the extensive vasculature that permeates the brain, and the potential for an NP to be within the vasculature (blood), associated with components of the blood–brain barrier (BBB) or blood–cerebrospinal fluid barrier (BCSFB), or in cerebrospinal fluid (CSF), rather than in brain parenchyma that houses the brain cells and their extracellular (interstitial) space. This article aims: to inform about the distinction between distribution into the brain (bulk brain) versus brain parenchyma, present methods that can demonstrate brain parenchymal distribution, present a tiered classification of methods and results that demonstrate bulk brain versus brain parenchymal distribution, critically review some reports that claimed brain or brain parenchymal NP distribution and rate them according to the tiered classification, and suggest that NP residence time in the brain often informs about brain parenchymal entry.

BBB anatomy & function

To justify the claim that an NP crossed the BBB and entered brain parenchyma requires an understanding of brain anatomy, the vasculature within it and research methods that demonstrate brain parenchymal entry. The primary route to enable NP brain parenchymal entry is across the BBB. The BBB separates the vascular compartment from brain cells that are surrounded by extracellular/interstitial fluid that occupies approximately 20% of the parenchymal space. The BBB includes brain microvascular endothelial cells (BMECs) that line microvessels within the brain, forming the luminal (blood) side of the BBB (Figure 1). BMECs have efflux transporters that effectively prevent brain parenchymal entry of many substances and metabolic enzymes to maintain brain homeostasis. On the other hand, BMECs have influx transporters that deliver essential nutrients, elements and factors to brain cells from the blood, supporting brain nutrition. Influx transporters have been utilized in a Trojan horse approach to hitchhike NPs through the BBB to brain parenchyma. BMECs differ from endothelial cells that line the vasculature outside of the brain by their near total absence of fenestrations (windows) through which NPs might diffuse and the presence
of tight junctions between adjacent BMEC surfaces. Intact tight junctions prevent passage of ionic substances as small as lanthanum (hydrated ionic diameter \( \sim 0.8 \text{ nm} \)) through this paracellular barrier [1]. Microvessels, the blood vessels between arteries and veins, are approximately 7.5 \( \mu \text{m} \) in diameter [2], barely large enough for a red blood cell to squeeze through. Microvessels deliver blood to the brain from within the brain [3].

There are 100 billion capillaries, approximately 600 km long, with a surface area of 20 m\(^2\) within the human brain. These microvessels provide a blood supply within approximately 10 \( \mu \text{m} \) of every brain cell [4], as illustrated in Figure 1 by the foreground and two background microvessels. They occupy approximately 2% of cortical brain volume and a greater space in some other brain regions [5,6]. Pericytes extend over approximately 30% of the BMEC surface. Surrounding the BMECs and pericytes is a basement membrane, covering the abluminal (brain-side) BMEC surface (Figure 1). Foot processes of astrocytes, one of the brain’s glial cells, cover more than 90% of the abluminal basement membrane surface (Figure 1). Neurons interact with this cell complex to form the neurovascular unit (Figure 1). For an NP to enter brain parenchyma from blood, it must either distribute through a BMEC (transcellular) or between BMECs (paracellular), as suggested in Figure 1. The latter is not expected when the BBB is intact but can occur when the BBB is opened (leaky, e.g., there is increased space between adjoining BMEC surfaces). This occurs in cancer (at the blood–tumor barrier), Alzheimer’s disease, traumatic brain injury, multiple sclerosis, infections, encephalitis, stroke, liver failure and other less common conditions [7,8]. BBB opening can be produced by osmotic insult, for example, 25% mannitol infusion into a carotid artery that delivers blood directly to its ipsilateral hemisphere [9] or focused ultrasound in conjunction with systemically circulating microbubbles [10].

The potential pitfall of experimental result misinterpretation

Given the intimate spatial relationship between the microvessels of the vascular compartment and brain parenchyma, visual demonstration of brain parenchymal entry \textit{in vivo} or \textit{in situ} requires \( \leq 10 \mu \text{m} \) resolution. Alternatively, one could use methods that separate BBB components from brain parenchymal components. Failure to distinguish brain parenchymal NP entry from bulk brain can result in the interpretation that an NP has crossed the BBB when it may not have or did not to the extent implied. For example, IVIS demonstration of increased fluorescence in the head region from an NP component does not differentiate whether the NP is in the skull; the blood in
the vessels surrounding and within the brain; associated with the luminal side of BMECs; within or between the BMECs; between the BMECs and basement membrane; within the basement membrane, pericytes or astrocyte foot processes; or in brain parenchyma.

There have been many studies addressing brain entry of many different NPs that concluded they entered the brain, without demonstration that they crossed the BBB and entered brain parenchyma. Considering the brain as the cranial contents, the claim of NP brain entry can be accepted if it is shown that the NP is in bulk brain. But this does not demonstrate brain parenchymal entry. For most studies, the methods employed were not able to differentiate NPs in the blood versus associated with BBB components versus residing in brain parenchyma, as differentiated in Figure 1.

NPs in or associated with the vascular compartment contribute to bulk brain but not brain parenchymal content

Rats perfused to remove blood (from the vascular compartment) 4 h after intravenous injection of gold glyconanoparticles had only approximately 4% as much NP component in bulk brain as rats that had not been perfused [11]. Similar perfusion reduced gold in three bulk brain regions to 7–18% of that seen in nonperfused rats [12]. This demonstrates that a great percentage of NPs in bulk brain may not be in brain parenchyma. Removal of blood from the entire body can be accomplished in the deeply anesthetized rodent by transcardial perfusion; introducing a perfusate into the left ventricle and ligating the right auricle to allow blood and perfusate drainage. Perfusate can be 0.9% sodium chloride or 0.1 mM phosphate @ pH 7.4. For electron microscopy, 4% paraformaldehyde and 3.5% glutaraldehyde in cacodylate/phosphate-buffered saline/saline buffer should be used. Gage et al. provide a protocol [13]. Some studies accounted for the contribution of blood NPs within the brain to their bulk brain level [6,14–18]. NPs in bulk brain blood were estimated from the product of the peripheral blood NP concentration × brain vascular volume. However, this does not account for NPs adsorbed to the BMEC luminal wall or in BBB cellular and membrane components. After intravenous injection to rodents, NPs were observed adhered to the luminal wall of blood vessels in the brain with little to no evidence of brain tissue entry. This was seen for lipid (stearic acid and polysorbate 80) drug conjugate NPs [19], 100-nm wide and 300-nm long poly(ethylene glycol) (PEG)-polyethylenimine-conjugated mesoporous silica NPs [20], and a 5-nm polyhedral citrate-coated ceria NP that had a surface charge of -53 mV [21]. In Dan et al.’s study, blood was removed from the brain by ceria-free perfusion. The preponderance of NPs in guinea pig brain that had been infused for 120 min with approximately 8-nm amphiphilic hydrophilic/hydrophobic-coated gold/iron core NPs followed by 30 min perfusate washout was in endothelial cells [22]. These observations are in agreement with a kinetic study that reported a negatively charged NP associated with cell surfaces within seconds, by Langmuir adsorption through electrostatic interaction [23]. These results demonstrate the potential contribution of BBB-associated NPs to bulk brain content, even after removal of blood from the brain.

The importance of distinguishing among NP distribution in the vascular compartment, the cells comprising the BBB and brain parenchyma is nicely illustrated by the results of Fiandra et al. [24]. Employing fluorometric dye fluorescein isothiocyanate (FITC)-labeled amphiphilic polymer-coated iron oxide NPs containing a fluorometric dye (AF660)-labeled antiretroviral drug, they differentiated localization of the drug from the nanconstruct. Confocal laser scanning microscopy (CLSM) viewing of mouse brain sections stained to identify the BMECs demonstrated colocalization of the NPs and BMECs, interpreted as NP residence in the vascular compartment. Antiretroviral drug was seen within and outside of the vascular compartment, interpreted as having been released from the NPs. Similarly, using a GFAP antibody to identify blood vessels by the astrocyte foot processes that cover them, Dal Magro et al. saw considerable NPs surrounded by the GFAP antibody, but some particles that were not, the latter interpreted as having entered brain parenchyma [25].

BCSF anatomy & function

The CSF compartment is comprised of four ventricles within the brain (two lateral, a third and a fourth) and the CSF-filled subarachnoid space that surrounds the entire CNS. CSF is produced by choroid plexuses in each of the four ventricles and diffusion of extracellular fluid from the brain. Each lateral ventricle has a single outlet, to the third ventricle, that drains into the fourth ventricle, that drains into the subarachnoid space, creating one-directional flow out of the brain. CSF in the human and rat turns over approximately four- and ten-times daily, respectively. It exits from this compartment through arachnoid granulations into dural sinuses then into blood (via the jugular veins). The epithelial cells of the choroid plexuses have tight junctions, efflux transporters and metabolic properties
similar to the BBB, creating the BCSFB between blood and CSF. There is little barrier to distribution between CSF and the brain, so substances injected into the CSF compartment, or able to cross the BCSFB, readily diffuse into the brain proximal to the CSF compartment.

Material entry into the nervous system, including the brain, might also be achieved by its introduction into the CSF compartment. In the human, this is most often accomplished by injection into CSF in the spinal region. Given CSF flow, NP injection into CSF would not be expected to persist for a long time compared with introduction into brain parenchymal space. As distribution from CSF into the CNS (brain and spinal cord) is by diffusion, and against extracellular fluid flow toward CSF, NP injection into CSF has been found to persist for much longer (Table 1 & Supplementary Figure 1).

**Convection-enhanced delivery of solutions into brain extracellular space**

A method to directly deliver solutions into the brain is convection-enhanced delivery (CED). In CED, the tip of a catheter is stereotaxically inserted into the brain through a hole in the cranium. A solute (drug or NP)-containing solution is infused through the catheter by a pump. Solute release from the catheter tip into brain extracellular space displaces extracellular fluid. The solute distributes through brain extracellular space surrounding the catheter tip by bulk flow to create a radial distribution up to a few centimeters. After termination of the infusion, the solute may continue to redistribute by bulk flow and diffusion as it is being locally cleared. CED has advantages of bypassing the BBB enabling brain parenchymal delivery of drugs that do not cross the BBB, delivery to a target site that can be deep within the brain and creation of much higher regional than distal brain and systemic drug concentration. It is most commonly used to treat glioblastoma multiforme, a brain cancer that has a poor prognosis [26–28]. Small molecules are cleared within 72 h [29], whereas some NPs have been found to persist for much longer (Table 1 & Supplementary Figure 1).

### Table 1. Studies that reported the brain level of the nanoparticle or component(s) at multiple times after convection enhanced delivery. Mean residence time values were determined as described in nanoparticle kinetics in the brain.

<table>
<thead>
<tr>
<th>Study summary</th>
<th>MRT (h)</th>
<th>Ref.</th>
</tr>
</thead>
<tbody>
<tr>
<td>The percent of radioactivity in the brain following CED of 82 nm 1H-cholesteryl hexadecyl ether liposomes was determined</td>
<td>22</td>
<td>[83]</td>
</tr>
<tr>
<td>The rat brain hemisphere that received CED of 0.06, 0.8 or 1.6 mg of 96–101 nm irinotecan-loaded nanoliposomes was perfused and homogenized, centrifuged, and irinotecan analyzed by the HPLC. Half-life of the three doses was 6.7, 10.7 and 19.7 days, respectively</td>
<td>108, 181 and 400</td>
<td>[81]</td>
</tr>
<tr>
<td>Brains from rats that received CED of 105–122 nm topotecan-loaded liposomes was perfused, the infused hemisphere removed, homogenized, centrifuged and topotecan analyzed by the HPLC. Half-life was 1.5 days</td>
<td>44</td>
<td>[84]</td>
</tr>
<tr>
<td>The rat brain hemisphere that received CED of a mixture of 95–110 nm doxorubicin-loaded pegylated liposomes or irinotecan-loaded nanoliposomes was perfused, homogenized, centrifuged and doxorubicin analyzed by the HPLC. Half-life was 10.9 and 16.7 days, respectively</td>
<td>273 and 305</td>
<td>[79]</td>
</tr>
<tr>
<td>Local retention of radioactivity after intratumor CED of 135 nm 186Re-loaded liposomes was determined in rat brain by gamma camera. Half-clearance time was 54.6 h</td>
<td>50</td>
<td>[85]</td>
</tr>
<tr>
<td>Brain MRI was obtained after CED of human serum albumin-, PEG- and methotrexate-human serum albumin-coated 20 nm γ-Fe2O3 NPs</td>
<td>190, 184 and 198</td>
<td>[80]</td>
</tr>
<tr>
<td>IVIS imaging was conducted after CED of EGFR-coated quantum dot-containing immunoliposome hybrid NPs to mice with brain tumors</td>
<td>97</td>
<td>[86]</td>
</tr>
<tr>
<td>Positron emission tomography imaging was conducted after CED of 71 and 147 nm [18F]-fluorobenzylamine, NHS-PEG-biotinylated, avidin-modified PLGA NPs to rats. Half-life was 111 min</td>
<td>0.39</td>
<td>[87]</td>
</tr>
<tr>
<td>Rat brain containing the region of CED of 175 nm carboxatin-labeled fluorescein-PLGA NPs was homogenized and platinum determined by ICP–MS</td>
<td>18</td>
<td>[88]</td>
</tr>
<tr>
<td>The hemisphere that received CED of 60–100 nm radiosensitizer (VE822)-loaded PEG-conjugated, poly(u-pentadecalactone-co-p-dioxanone) NPs, of varying PEG polymer ratios, was analyzed by LC–MS for VE822. Half-life was 12 h, 2 and 5 days</td>
<td>46, 61 and 68</td>
<td>[89]</td>
</tr>
<tr>
<td>Fluorescence images of mice brain sections were obtained after intrastratial CED of Cy5.5 fluorophore-labeled peptide-based nanofibers (5 nm width, 100, 400 and 1000 nm length)</td>
<td>20, 25 and 29</td>
<td>[29]</td>
</tr>
<tr>
<td>MRI imaging was conducted for 48 h after CED of 54 nm iron oxide chitosan-PEG NPs modified with a glioma-targeting ligand to mice-bearing brain tumors. Signals near the point source of the infusion of NP without and with ligand were used to calculate the MRT</td>
<td>3.4 and 4.8</td>
<td>[90]</td>
</tr>
</tbody>
</table>

CED: Convection-enhanced delivery; HPLC: High-performance liquid chromatography; ICP–MS: Inductively coupled plasma mass spectrometry; IVIS: In vivo imaging system; LC–MS: Liquid chromatography-mass spectrometry; MRT: Mean residence time to the last sample; NHS: N-hydroxysuccinimide; NP: Nanoparticle; PEG: Polyethylene glycol; PLGA: Poly(D,L-lactide-co-glycolide).
Table 2. Visual methods that might be used to image brain nanoparticle distribution and their reported resolution.

<table>
<thead>
<tr>
<th>Method</th>
<th>Resolution</th>
<th>Ref.</th>
</tr>
</thead>
<tbody>
<tr>
<td>High-resolution brain MRI</td>
<td>$\geq 120 \ \mu m$</td>
<td>[91]</td>
</tr>
<tr>
<td>Photoacoustic microscopy</td>
<td>20–100 $\mu m$</td>
<td>[37]</td>
</tr>
<tr>
<td>IVIS spectrum</td>
<td>20 $\mu m$</td>
<td>[92]</td>
</tr>
<tr>
<td>MSI</td>
<td>Submicron for some methods</td>
<td>[93]</td>
</tr>
<tr>
<td>Super resolution (light) microscopy (aka: single-molecule localization microscopy)</td>
<td>&lt;100 nm</td>
<td>[38,94–96]</td>
</tr>
<tr>
<td>Hyperspectral imaging</td>
<td>&lt;10 nm</td>
<td>[44,97,98]</td>
</tr>
<tr>
<td>CLSM</td>
<td>2 $\mu m$</td>
<td>[99]</td>
</tr>
<tr>
<td>Electron microscopy</td>
<td>&lt;1 nm</td>
<td>[100]</td>
</tr>
<tr>
<td>CLEM</td>
<td>&gt;50 nm for LM, &lt;1 nm for EM</td>
<td>[59,61]</td>
</tr>
</tbody>
</table>

CLEM: Correlative light and electron microscopy; CLSM: Confocal laser scanning microscopy; EM: Electron microscopy; IVIS: In vivo imaging system; LM: Light microscopy; MSI: Mass spectrometry imaging.

The nasal cavity route of delivery directly to the CNS

There is only one site where the nervous system is directly exposed to the external environment of mammals, where direct uptake of NPs into the nervous system might occur. This is the roof of the nasal cavity where the olfactory neuron and the maxillary branch of the trigeminal neuron have terminals with receptors that mediate the perception of sensory stimuli, including smell. This route of administration, introducing test material into the nose (intranasal), is being investigated for NP entry into the brain, to bypass the BBB. Demonstration of NPs in these sensory neurons ex vivo (e.g., by electron microscopy visualization or elemental analysis under conditions that rule out dissolution or degradation that releases NP components), or of NPs within the nerve when it is in situ, provides confidence of nerve entry. The olfactory and trigeminal neurons are surrounded by CSF, so suggestion of NP in the region of the nerve, for example, from an NP fluorescent molecule, does not unequivocally show nerve entry. Although neuropeptides have been shown to enter CSF directly from the nasal cavity, this route was shown to not contribute to uptake of 110 nm 6-coumarin-loaded (for visualization) PEG-poly(lactic acid) NPs [30]. There have been demonstrations of NP uptake by the olfactory or trigeminal nerve. NPs often accumulate in the olfactory bulb, with little or no distribution across the synapse between the olfactory bulb and the next neuron of the olfactory system that would enable distribution to distal brain regions (e.g., the pyriform cortex, amygdala, thalamic and hypothalamic nuclei, and hippocampus) after intranasal administration. Olfactory bulb accumulation was shown with colloidal gold [31,32], $^{13}$C carbon [33], nanoscale manganese oxide [34] and quantum dots [35]. Distribution of quantum dots from the nasal cavity to the olfactory bulb was attributed to microtubule-mediated fast axonal transport (200–400 mm/day), because axonal flow is too slow to account for the timecourse of its appearance in the olfactory bulb [35]. Based on the olfactory nerve length from the nasal cavity to the olfactory bulb (~5 and 8 mm in mice and rats, respectively) fast axonal transport could traverse this with an NP in approximately 26–40 min. Much more time would be required for neuronal NP transport from the nasal cavity to distant brain regions, such as the striatum. Rapid appearance of NPs or NP components in distant sites can be attributed to absorption from the nasal cavity into systemic circulation followed by distribution into the brain. Translocation of NPs from the nasal cavity to the brain is low. Gamma counting of whole brain 24 h after intranasal $^{111}$In-labeled protein NP suggested approximately 0.2% of the dose was in the brain, assuming stability of the $^{111}$In-NP complex [36]. No studies were found demonstrating NP entry into brain parenchyma beyond the olfactory bulbs after intranasal administration.

Methods to demonstrate NP brain parenchymal entry

To definitively claim that an NP has entered brain parenchyma requires methods that differentiate NP distribution in brain parenchyma from blood within the brain and BBB components. One approach is to use visual methods.

Visual methods to demonstrate NP brain parenchymal entry

Table 2 lists some visual methods and their optimal resolution. High-resolution brain MRI does not provide, even under optimal conditions, sufficient resolution to differentiate NP localization between parenchymal versus nonparenchymal sites. It would not be able to characterize NP morphology, size or chemical identity.

Photoacoustic imaging (tomography) is a hybrid imaging modality that integrates optical contrast with high-ultrasonic spatial resolution in deep tissue. It can provide noninvasive images of the entire brain with less than msec
temporal resolution [37], based on the light absorption properties of the tissue and material (e.g., NPs) within it. Although useful for NP-assisted imaging, for example, of brain tumors, it lacks sufficient resolution to differentiate NP localization between parenchymal versus nonparenchymal sites.

The IVIS spectrum in vivo imaging system combines 2D and 3D optical tomography into a single platform to visualize fluorescence and bioluminescence. It has been extensively used to demonstrate the intensity of fluorescent NPs in organs in vivo and ex vivo. Intensity in the head region has been claimed by many as evidence of the NP in the brain. It is noninvasive and can be used with living organisms. However, even if used to localize NPs in ex vivo tissue, it does not have sufficient resolution to differentiate NP localization between parenchymal versus nonparenchymal sites.

Mass spectrometry imaging (MSI) includes many techniques, such as laser desorption/ionization mass spectrometry, matrix-assisted laser desorption ionization, (time-of-flight) secondary ion mass (SIM) spectrometry and desorption electrospray ionization. It has been used with surface-functionalized gold- or gadolinium-containing NPs to localize organics such as lipids, metabolites and proteins in the brain. Although some forms of MSI have submicron resolution (e.g., SIM), no reports were found using MSI to verify brain parenchymal NP localization.

Super-resolution microscopy utilizes fluorescent microscopes in a variety of approaches, categorized as those that approach the diffraction limit (0.25 μm in the focal plane) and those that break the diffraction limit by turning fluorescent markers on and off. In the latter rather than all fluorescent molecules emitting simultaneously (as occurs in conventional fluorescence microscopy), a small subpopulation is excited so that emission from neighbors does not overlap, enabling isolation of individual emitters [38]. Approaches include stochastic optical reconstruction microscopy, direct stochastic optical reconstruction microscopy, photoactivated localization microscopy, fluorescence photoactivated localization microscopy and ground-state depletion with individual molecule return. Super-resolution microscopy has been used to image the NP protein corona [39,40] and localize NPs in cells [41,42], but no reports were found of in vivo, in situ or ex vivo NP localization.

Hyperspectral imaging combines imaging and spectrophotometry by capturing spectral data of infrared and visible light reflected by samples at each pixel in an enhanced dark-field microscopic image. For this to differentiate vascular versus parenchymal NP localization, an enhanced dark-field microscopic image needs to be interpreted first to identify the region of interest, then hyperspectral imaging employed to provide spectrophotometric data. This was demonstrated in studies that showed cellular sites of functionalized gold NPs [43]; studies that included cellular, C. elegans, and Japanese medaka NP distribution cited by Roth [44]; and porcine skin uptake of metal oxide NPs [45]. This approach might enable differentiation of NP localization between brain parenchymal versus vascular sites, but no such studies were found.

Fluorescence microscopy visualizes fluorescent tags or incorporated molecules (e.g., doxorubicin and paclitaxel). With the exception of transmission electron microscopy (TEM), the only studies using a method in Table 2 that appear to have differentiated NP parenchymal from vascular space distribution used fluorescence microscopy and CLSM. Using fluorescence microscopy, Mulik et al. saw fluorescent-labeled NPs within brain cells [46]. Åsund et al. used CLSM to image fluorescent dye-containing NPs, and tomato lectin-labeled blood vessels, and saw NP fluorescent dye outside of the brain microvasculature [47]. Some details of these studies and those reported by others who used CLSM are below and/or in Table 4 [22,24,25,47–55].

Electron microscopy, typically TEM, is not only the best technique to characterize prepared NPs, it is also amenable to physicochemically identify and characterize NPs in samples obtained from biological environments [56–58]. It provides the greatest resolution among the options in Table 2, and when equipped with capabilities such as high-angle annular dark-field detection, energy dispersive x-ray spectroscopy or electron energy loss spectroscopy, it enables chemical characterization.

The benefits of light microscopy (including identification of biological structures and location of regions of interest and fluorescent labels) have been combined with the benefits of electron microscopy (to provide NP localization and physicochemical information) in correlative light and electron microscopy (CLEM) [59]. The two approaches were independently used to locate NPs in a murine brain tumor model [60]. CLEM has been developed in a single instrument and used to localize NPs in cultured non-neuronal cells [61]. No studies were found utilizing CLEM to investigate NP brain parenchymal entry.

Several of the above methods are based on detection of a label added to the NP, often a fluorescent molecule or metal/metal oxide. Correct interpretation of NP localization, when based on an added label, requires NP preparation that does not contain free label (which can be removed by dialysis) and understanding the label stability in vivo. Fluorescent molecules are susceptible to loss of their signal due to quenching (due to interaction with local molecular
environmental components) and photobleaching (excitation light-induced destruction of the excited fluorophore). Persistent association (stability) of the label and the NP in vivo is required to attribute label signal to NP localization. Label dissociated from the NP will yield a different distribution than the label-NP complex. Covalent binding of label to an NP decreased label release, compared with encapsulation of the label [62]. Incorporation of the label in the NP core did not produce changes in the NP surface properties [63]. Label dissociation from a biodegradable polymer NP would be expected over time. Label dissociation can be assessed in vitro by incubation of the labeled NP with brain homogenate [30] or separation of the NP and brain components from the released label, for example, by NP centrifugation, and detection of label in the supernatant. Other considerations include the potential for the label to alter the physicochemical properties of the NP and its resultant distribution and biological tolerance to the label. More in-depth discussions can be found in [64,65].

**Procedural methods to demonstrate NP brain parenchymal entry**

Methods are available to assess whether NPs introduced into the vasculature perfusing the brain are associated with BBB components and/or crossed the BBB to enter brain parenchyma. The in situ brain perfusion technique can be used to determine NP entry rate and extent of distribution into bulk brain, multiple bulk brain regions and/or the choroid plexus, ipsilateral to the carotid artery perfused after short-term (few minutes or less) intracarotid infusion [66]. An intracarotid perfusion rate sufficiently high to prevent blood from entering the perfused carotid artery is used, therefore preventing blood from entering the brain hemisphere perfused by that artery. This enables control of the chemical environment of the material tested (therefore its chemical form [speciation]), based on the perfusion fluid composition and avoidance of blood exposure. Preventing blood exposure avoids potential NP biotransformation (e.g., corona coating by plasma proteins, dissolution or particle breakdown) that might change its surface properties and brain uptake. Although the in situ brain perfusion technique removes blood from the brain vasculature, it does not distinguish NP distribution associated with the BBB versus brain parenchyma, nor choroid plexuses from brain unless they are separated.

The capillary depletion method is intended to separate BBB components from brain parenchyma, to produce capillary-depleted brain parenchyma [67]. This is often conducted with brain tissue obtained using the in situ brain perfusion technique but can be used with brain tissue obtained by other methods that remove blood from the brain. The methods are described in detail in [68]. The capillary depletion method uses centrifugation of brain homogenate in dextran that separates the BMECs into the pellet (along with brain nuclei and erythrocytes if blood was not perfused from the brain) from the supernatant that contains brain cells and brain extracellular fluid. Presence of the test material in the supernatant is taken as evidence that it has transcytosed the BBB into brain parenchyma. This method was employed to ascertain the fraction of 190-nm poly(n-buty 1cyanoacrylate) (PBCA)-coated doxorubicin-loaded NPs that entered brain parenchyma [69]. Doxorubicin was seen in the brain fraction after intravenous administration in NPs but not when doxorubicin solution was administered with 1% polysorbate 80 (intended to open the BBB). Coating the NPs with 1% polysorbate 80 increased the doxorubicin supernatant to pellet ratio, suggesting the surfactant-coated PBCA NPs entered brain parenchyma. Employing the capillary depletion method after whole body perfusion to remove blood, it was shown that the percentage of the injected dose of [111In-DTPA-multiwalled carbon nanotubes (median diameter: 18.9 and length: 500 nm) decreased in brain capillaries but not brain parenchyma over 24 h [70]. The capillary depletion method was used to obtain curcumin for LC–MS/MS analysis from mouse brains after oral dosing of a curcumin-containing NP. The report does not state if the brain was perfused prior to harvest. If not the supernatant fraction would be expected to include blood plasma contents that could have contained some/all of the curcumin [71]. The capillary depletion method was used to demonstrate that cationic bovine serum albumin-coated biodegradable polymersomes (∼100 nm, composed of PEG- and maleimide-PEG-poly(ε-caprolactone), loaded with coumarin-6) entered brain parenchyma. The supernatant fraction increased from a half to 1 h and was fairly stable for 4 h, and increased over the 4 h in relation to the pellet (vascular fraction) [72]. Capillary depletion was used in transcardially perfused mice to assess the influence of transferrin-receptor antibody density on gold NPs and platinum-containing liposomes to enter brain parenchyma [73]. 2.5 h after intravenous injection, gold and platinum in the parenchymal fraction were approximately 27 and 22% of the combined parenchymal and capillary fractions, respectively. In contrast to demonstrating brain parenchymal entry, the capillary depletion method suggested nearly all of 5-nm ceria in the brain was associated with the capillary endothelial cells, consistent with EM observations of nanoceria adhered to the luminal side of blood vessel walls in the brain, noted above [21].
NP components have been sampled from the brain using microdialysis probes [74–77]. In the presence of an intact BBB, it is assumed that this technique samples extracellular space. Microdialysis probes with 30 and 100 kDa molecular weight cutoff membranes would be expected to allow penetration of up to 5–8 and 8–14 nm (depending on membrane composition) diameter spherical particles and molecules released from NPs. The appearance of an NP component in the dialysate from a probe would not conclusively demonstrate NP distribution into brain parenchyma. None of the above studies demonstrated that intact NPs were recovered in the dialysate.

**Level of demonstration of NP brain or brain parenchymal entry & critical review of study claims**

Convincing demonstration of brain parenchymal entry is limited to in vivo studies. In vitro BBB models do not recapitulate BBB complexity. They lack all of its cell composition, cell interactions, cell spatial relationships and magnitude of resistance to distribution across them. Therefore, this report is limited to in vivo, in situ and ex vivo brain studies. Many reports published within the past 5 years were identified by a SciFinder search of the terms NP and brain, a SciFinder search of the terms NP and BBB for reports published in 2019, and a PubMed search of NPs, drug delivery systems and brain parenchyma. The reports were reviewed, summarized and critiqued for evidence supporting claims in the report and classified for the level of demonstration of NP brain or brain parenchymal entry, as defined in Table 3. Some reports published more than 5 years ago that are informative for their demonstration of NP brain parenchymal entry and/or reported brain NP concentration at multiple times are included. The results are in Table 4.

**NP kinetics in the brain**

Endocytosis is the prevalent mechanism that mediates NP transport across the intact BBB. It would not be anticipated that after NPs cross the BBB and enter brain parenchyma they would be rapidly cleared from that compartment [78]. After NP infusion directly into the brain by CED, HPLC determination of two components showed clearance half-lives of 250–400 h (Table 1) [79]. MRI-determined NP distribution volume after CED decreased 50% in approximately 250 h (Table 1) [80]. Some details of these and other studies that reported brain NP component levels at multiple times after CED are in Table 1. Brain clearance half-life was shown to be dose

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**Table 3. A classification of results demonstrating brain versus. brain parenchymal entry.**

<table>
<thead>
<tr>
<th>Level</th>
<th>Description of results</th>
<th>Interpretation</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>NP component (e.g., fluorescent dye, element) in bulk brain or ex vivo brain not perfused of blood. Determined by methods such as IVIS, radioactivity, NAA, ICP-MS, AAS or HPLC</td>
<td>A NP component was in bulk brain</td>
</tr>
<tr>
<td>2</td>
<td>NP component (e.g., fluorescent dye, element) in bulk brain or ex vivo brain perfused to remove blood. Determined by methods such as IVIS, radioactivity, NAA, ICP-MS, AAS or HPLC</td>
<td>A NP component was in bulk brain, but not in the blood within the brain</td>
</tr>
<tr>
<td>3</td>
<td>Intact NP in bulk brain not perfused of blood. Determined by methods such as bulk brain homogenization, NP isolation and verification (e.g., visual, perhaps also requiring chemical)</td>
<td>The NP was in bulk brain</td>
</tr>
<tr>
<td>4</td>
<td>Intact NP in bulk brain perfused to remove blood. Determined by methods such as bulk brain homogenization after perfusion to remove blood, NP isolation and verification (e.g., visual, perhaps also requiring chemical)</td>
<td>The NP was in bulk brain, but not in the blood within the brain</td>
</tr>
<tr>
<td>5</td>
<td>NP component isolated from brain parenchyma. For example, using the capillary depletion method of brain perfused to remove blood to obtain brain parenchyma and a method to identify the NP component, for example, radioactivity, ICP-MS, AAS or HPLC. For example, microdialysis to recover an NP component from brain extracellular fluid, in the presence of an intact BBB</td>
<td>A NP component was in brain parenchyma</td>
</tr>
<tr>
<td>6</td>
<td>NP component (e.g., fluorescent dye, element) visualized in brain areas or cells not associated with brain vasculature. Determined by methods such as TEM, with chemical verification, such as EDS or EELS</td>
<td>A NP component was in brain parenchyma</td>
</tr>
<tr>
<td>7</td>
<td>Intact NP isolated from brain parenchyma. For example, using the capillary depletion method of brain perfused to remove blood to obtain brain parenchyma and a method to verify intact NPs (e.g., visual, perhaps also requiring chemical)</td>
<td>The NP was in brain parenchyma</td>
</tr>
</tbody>
</table>

The level of demonstration of nanoparticle brain or brain parenchymal entry is in the first column.

Brain delivery

Whole organs were homogenized, proteins precipitated and curcumin determined by HPLC after iv. injection of 90-nm doxorubicin-loaded solid lipid particles

HPLC results of NP component show bulk brain uptake; 1

1.8 (1.8) [102]

Brain delivery

Whole organs were homogenized, proteins precipitated and doxorubicin determined by HPLC after iv. injection of 70–345 nm methotrexate-loaded PBCA NPs

HPLC analysis of NP component in bulk brain does not demonstrate distribution across the BBB; 1

1.8 and 1.9 [103]

‘Polysorbate 80-coated polybutylcyanoacrylate NPs could be used to overcome blood–brain barrier’

Visible blood vessels were removed from the cerebrum and cerebellum, whole organs were homogenized, proteins precipitated and methotrexate determined by HPLC after iv. injection of 17- and 37-nm gold particles.

ICP–MS results indicated that 17 and 37 nm (gold NPs) were capable of passing through the blood–brain barrier and entering the mouse brain.‘

ICP–MS results of NP component in bulk brain do not demonstrate distribution through the BBB; 1

Increased from 336 to 504 h after injection

Table 4. Brain nanoparticle distribution study claims, summary, critique, support for the claims and level of brain or brain parenchymal entry.

<table>
<thead>
<tr>
<th>Claim</th>
<th>Study summary</th>
<th>Critique, support for claim and level of demonstrated NP brain entry</th>
<th>MRT (h)</th>
<th>Ref.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Uptake of NPs in the brain</td>
<td>Whole organs were solubilized for radioactivity determination after iv. injection of 131-nm surfactant-coated poly[14C-methyl methacrylate] particles</td>
<td>Claim assumes intact NP distribution to the brain, which is not shown. Bulk brain 14C presence shown; 1</td>
<td>42 (47)</td>
<td>[101]</td>
</tr>
<tr>
<td>Doxorubicin was in brain tissue</td>
<td>Whole organs were homogenized, proteins precipitated and doxorubicin determined by HPLC after iv. injection of 90-nm doxorubicin-loaded solid lipid particles</td>
<td>HPLC results of NP component show bulk brain uptake; 1</td>
<td>1.8 (1.8)</td>
<td>[102]</td>
</tr>
<tr>
<td>‘Polysorbate 80-coated polybutylcyanoacrylate NPs could be used to overcome blood–brain barrier’</td>
<td>Visible blood vessels were removed from the cerebrum and cerebellum, whole organs were homogenized, proteins precipitated and methotrexate determined by HPLC after iv. injection of 17- and 37-nm gold particles</td>
<td>ICP–MS results of NP component in bulk brain does not demonstrate distribution across the BBB; 1</td>
<td>1.8 and 1.9</td>
<td>[103]</td>
</tr>
<tr>
<td>‘ICP–MS results indicated that 17 and 37 nm (gold NPs) were capable of passing through the blood–brain barrier and entering the mouse brain.’</td>
<td>‘Whole brain was homogenized, separated by HPLC, and gold quantified by ICP–MS after iv. injection of 17- and 37-nm gold particles. TEM and EDS showed electron-dense particles in the cytoplasm of hippocampal pyramidal cells and gold electron densities are iron; 6 TEM demonstrates delivery across the BBB; 1’</td>
<td>ICP–MS results of NP component in bulk brain do not demonstrate distribution through the BBB; 1; TEM/EDS results demonstrate distribution across the BBB; 6</td>
<td>1.8 (1.8)</td>
<td>[104]</td>
</tr>
<tr>
<td>Brain delivery</td>
<td>Whole brain from transversally perfused rats was homogenized for gold quantitation by NAA after iv. injection of peptide-conjugated, citrate-coated 12-nm gold particles. Whole brain from transversally perfused rats were prepared for fluorescence microscopy after iv. injection of carboxyfluorescein-labeled, peptide-conjugated, citrate-coated 12-nm gold particles</td>
<td>NAA results of NP component in brain homogenate and concurrent fluorescence show bulk brain uptake; 2</td>
<td>No consistent increase or decrease over 24 h</td>
<td>[105]</td>
</tr>
<tr>
<td>‘Focused ultrasound and magnetic targeting synergistically delivers therapeutic MNPs across the blood–brain barrier to enter the brain’</td>
<td>‘Whole brain was prepared for ICP–OES iron and HPLC epirubicin determination after iv. injection of 12-nm magnetic NPs into rats whose BBB had been opened by iv. injection of microbubbles and focused ultrasound. TEM of similarly treated rats with brain tumors showed electron densities inside tumor cells and macrophages</td>
<td>ICP–OES and HPLC demonstrate bulk brain but not parenchymal entry; 1</td>
<td>11 [106]</td>
<td></td>
</tr>
<tr>
<td>‘NP deposits were widely distributed throughout brain, with apparent cellular uptake’</td>
<td>‘Microscopic assessment of sections from rat brain after intranasal dosing of 299-nm fluorescent- and ovalbumin-loaded cationic liposomes revealed the label in the striatum and substantia nigra. TEM and EDS showed electron-dense particles in the cytoplasm of hippocampal pyramidal cells and gold electron densities are iron; 6 Imaging shows increased fluorescence within cells; 5 Colocalization shows fluorescence from NP within cells; 5</td>
<td>Image shows increased fluorescence within cells; 5</td>
<td>4.3 and 2.1 [36]</td>
<td></td>
</tr>
<tr>
<td>‘Curcumin-loaded PLGA NPs could cross the blood–brain barrier (BBB) to enter brain tissue’</td>
<td>Brains from perfused rats were dissected into parts, homogenized, proteins precipitated and curcumin determined by HPLC after iv. injection of 163-nm curcumin-loaded PLGA NPs. Reported MRT was 27 min</td>
<td>HPLC analysis of NP component in bulk brain does not demonstrate distribution across the BBB; 2</td>
<td>0.36 (0.33)</td>
<td>[107]</td>
</tr>
<tr>
<td>‘In vivo brain distribution … monitored by a real-time fluorescence imaging system’</td>
<td>The whole rat and ex vivo brain were bioluminescence imaged using IVIS and fluorescence microscopic images obtained of brain sections after iv. injection of 190–210 nm fluorescent dye-labeled solid lipid particles</td>
<td>Imaging demonstrates bulk brain distribution; 1</td>
<td>0.45 [108]</td>
<td></td>
</tr>
<tr>
<td>Brain delivery</td>
<td>Fluorescence intensity was determined in ex vivo organs from perfused rats after intranasal dosing of ovalbumin-conjugated fluorescent-labeled 90-nm PEG-PLGA NPs</td>
<td>Fluorescence intensity of brain demonstrates bulk brain delivery; 2</td>
<td>5.2 [109]</td>
<td></td>
</tr>
</tbody>
</table>

Mean residence time from studies that reported brain and blood (in parentheses) levels at multiple times after dosing. Study multiple end points are designated by a, b, c before the study summary and critique. The level of demonstrated nanoparticle brain entry is from the classification described in Table 3. The numbers after the sentences in column 3 refer to the level of demonstrated NP brain entry. AAS: Atomic absorption spectroscopy; BBB: Blood–brain barrier; BTB: Blood–brain tumor barrier; BMEC: Brain microvascular endothelial cell; BSA: Bovine serum albumin; CLSM: Confocal laser scanning microscopy; EDS: Energy dispersive x-ray spectroscopy; EELS: Electron energy loss spectroscopy; IRTC: Fluorescein isothiocyanate; ICP–MS: Inductively coupled plasma–mass spectrometry; ICP–OES: Inductively coupled plasma–optical emission spectroscopy; INAA: Instrumental neutron activation analysis; ip.: Intraperitoneal; iv.: Intravenous; IVIS: In vivo imaging system; MRT: Mean residence time; MS: Mass spectrometry imaging; NAA: Neutron activation analysis; NP: Nanoparticle; PBCA: Poly(14C-methyl methacrylate); RVG: Rabies virus glycoprotein; TEM: Transmission electron microscopy.
### Table 4. Brain nanoparticle distribution study claims, summary, critique, support for the claims and level of brain or brain parenchymal entry. (cont.)

<table>
<thead>
<tr>
<th>Claim</th>
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<th>Ref.</th>
</tr>
</thead>
<tbody>
<tr>
<td>‘We observed . . . a fourfold increase of silver content . . . which indicates that AgNPs are able to cross the blood–brain barrier (BBB)’</td>
<td><em>Brain from perfused rats was digested and silver determined by ICP–MS after iv. 20- and 200-nm silver NP injection.</em>&lt;br&gt;<strong>TEM showed electron-dense particles and clusters of particles in brain cells. Their elemental composition was not reported</strong></td>
<td><em>ICP–MS results of NP component in bulk brain do not demonstrate distribution across the BBB; 1.</em>&lt;br&gt;*<em>NPs crossed the BBB if the electron dense particles contained silver that was not the result of dissolution then precipitation; 6.</em></td>
<td>Increased from 24 to 672 h after injection</td>
<td>[110]</td>
</tr>
<tr>
<td>‘AgNPs also gained access to the brain . . . visible as single NPs as well as clusters’</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>‘NPs were localized in the brain parenchyma’</td>
<td><em>Gold in brains from perfused rats was determined by INAA after ip. injection of transferrin receptor targeting and β-amylid destroying peptides conjugated to 37-nm gold NPs.</em>&lt;br&gt;<strong>TEM of hippocampus and cortex of perfused rats after ip. injection of these gold NPs demonstrated intracellular electron dense spots</strong></td>
<td><em>INAA results of NP component in bulk brain do not demonstrate distribution across the BBB; 1.</em>&lt;br&gt;<strong>NP seen in brain parenchyma; 6,</strong>&lt;br&gt;<strong>assuming the spots were gold NPs</strong></td>
<td>8.8</td>
<td>[111]</td>
</tr>
<tr>
<td>‘Suggest that the OX-26-coated NPs were taken up by the brain’</td>
<td>Microdialysis samples were obtained from the ventral hippocampus for fluorescence detection of FITC-dextran after iv. administration of 140–160 nm CD71/OX26-coated FITC-dextran loaded methoxy-PEG-poly(lactic acid) NPs</td>
<td>NPs were too large to diffuse through the microdialysis membrane suggesting FITC-dextran release; 5, assuming intact BBB</td>
<td>21</td>
<td>[74]</td>
</tr>
<tr>
<td>‘Brain endothelial uptake and possibly parenchymal penetrability’</td>
<td>CLSM of fixed brain sections after carotid artery (in situ) brain perfusion and washout of perfusate or iv. injection of fluorescent dye-labeled, polyethylenimine/oligonucleotide-containing liposomes showed fluorescence associated with BMECs and in the perivascular space between the BMEC membrane and parenchymal basement membrane</td>
<td>NP or a component seen beyond BMECs but not in brain parenchyma; 2</td>
<td></td>
<td>[51]</td>
</tr>
<tr>
<td>‘NPs possess an appreciable BBB penetrability’</td>
<td>Microdialysis samples were obtained from the right ventrolateral periaqueductal gray for fluorescence detection of FITC-neurotoxin after intranasal dosing of 145-nm chitosan-modified PLA FITC-neurotoxin-containing NPs</td>
<td>NPs were too large to diffuse through microdialysis membrane suggesting FITC-neurotoxin release; 5, assuming intact BBB</td>
<td>3.3</td>
<td>[75]</td>
</tr>
<tr>
<td>‘Accumulation was achieved in the brain’</td>
<td>Whole brains from mice were homogenized, proteins precipitated, and camptothecin determined by HPLC after iv. injection of 160 nm camptothecin-containing solid lipid NPs.</td>
<td>HPLC analysis demonstrates bulk brain accumulation; 1</td>
<td>7.7(7.0)</td>
<td>[112]</td>
</tr>
<tr>
<td>‘[Palmitoyl dalargin] was detectable . . . in the brain’ <em>pDal is clearly located in the brain parenchyma</em></td>
<td><em>Whole brains were homogenized, proteins precipitated, and palmitoyl dalargin measured by LC–MS in rats after iv. injection of palmitoyl dalargin fibers. The authors claim brain parenchymal entry because brain concentration was greater than can be accounted for by blood, and slower clearance from brain than blood.</em>&lt;br&gt;<strong>Coherent anti-Stokes Raman scattering microscopy revealed pDal fibers within brain parenchyma</strong></td>
<td><em>LC–MS of NP component in bulk brain does not rule out BMEC adsorption or incorporation; 1.</em>&lt;br&gt;<strong>Microscopy demonstrated brain parenchymal fiber entry; 6</strong></td>
<td>0.8(0.17)</td>
<td>[113]</td>
</tr>
<tr>
<td>‘Ag detected in the brain and passing through the blood–brain barrier (i.e., not localized in the vessels and capillaries) constitutes no less than 93% of the total content of Ag in the organ’</td>
<td>Silver and gold were determined by NAA in brain of exsanguinated rats after 14 or 28 daily intragastric administrations of 35-nm silver and gold NPs. The authors calculated brain metal from its concentration in bulk and brain blood mass</td>
<td>This does not account for metal in BMECs. Silver could be ion released from NP dissolution. BBB Ag passage not shown; 2</td>
<td></td>
<td>[14]</td>
</tr>
<tr>
<td>‘In vivo studies indicated BBB penetration of intravenously injected TIMP-1 NPs + Pb80’</td>
<td>Confocal microscopy (40× objective) immunohistochemistry of brain slices from perfused mice after iv. injection of polysorbate 80-coated coumarin 6-containing protein-loaded PLGA NPs showed a few positive spots</td>
<td>Spot location (intra- vs extravascular) cannot be determined. BBB penetration not demonstrated; 2</td>
<td></td>
<td>[114]</td>
</tr>
</tbody>
</table>

Mean residence time from studies that reported brain and blood (in parentheses) levels at multiple times after dosing. Study multiple end points are designated by a, b, c before the study summary and critique. The level of demonstrated nanoparticle brain entry is from the classification described in Table 3. The numbers after the sentences in column 3 refer to the level of demonstrated NP brain entry.

Table 4. Brain nanoparticle distribution study claims, summary, critique, support for the claims and level of brain or brain parenchymal entry. (cont.).

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</tr>
</thead>
<tbody>
<tr>
<td>'BBB-penetrating cancer cell-targeting delivery system'</td>
<td>Whole animal and ex vivo brain IVIS imaging were conducted after iv. 85-nm fluorescent-labeled, doxorubicin-loaded desoxanoyl ghrelin and folate conjugated-polymerosome administration to brain tumor-bearing mice.</td>
<td>+Whole brain results using this method do not have sufficient resolution to demonstrate BBB penetration; 1.</td>
<td>[115]</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Brains from brain tumor-bearing perfused rats were homogenized, proteins precipitated and doxorubicin determined by HPLC after iv. injection of 85-nm fluorescent-labeled, doxorubicin-loaded desoxanoyl ghrelin and folate-conjugated polymersomes</td>
<td>+HPLC analysis of NP component in bulk brain does not demonstrate BBB penetration; 2</td>
<td></td>
<td></td>
</tr>
<tr>
<td>'Targeted ⁶⁸Ga-GNPs improves BBB crossing'</td>
<td>Whole brains from perfused rats were gamma counted after iv. injection of glucose conjugate-coated ⁶⁸Ga-gold NPs. Brain gold content was determined by ICP-MS</td>
<td>Whole brain radioactivity and ICP-MS results of NP component in bulk brain do not verify BBB crossing; 2</td>
<td>[11]</td>
<td></td>
</tr>
<tr>
<td>Brain delivery</td>
<td>Whole organs were homogenized and Bacoside-A determined by HPLC after ip. administration of 173-nm polyserabote 80-coated, Bacoside-A-loaded PLGA NPs to rats</td>
<td>HPLC analysis of NP component demonstrates bulk brain delivery; 1</td>
<td>[116]</td>
<td></td>
</tr>
<tr>
<td>'Brain bioavailability of brevicsapine from the BRE-GBSN-P5 and BRE-GBSN-P5-DSPE was significantly enhanced'</td>
<td>Fifteen KDa cutoff (∼ 5 nm) microdialysis probes were used to obtain dialysate from rat brain after iv. injection of 23-nm PEG, steartate, 1,2-dioeyl-sn-glycero-3-phophoethanolamine-N-PEG2000-coated brevicsapine-loaded solid lipid NPs. The primary active component of brevicsapine was measured by HPLC.</td>
<td>NP released component does not identify NP brain location; 5, assuming intact BBB</td>
<td>[76]</td>
<td></td>
</tr>
<tr>
<td>'We ... demonstrated the ability of these gelatin NPs to pass into the brain parenchyma following intranasal administration'</td>
<td>SEM of rat brain 6 h after intranasal administration of 184-nm gelatin NPs showed clusters of approximately 40-nm spherical particles in the olfactory tract, cortex, striatum, amygdala and hypothalamus as evidence of NP brain parenchymal entry</td>
<td>Assuming the observed particles were gelatin NPs, perhaps smaller due to tissue processing, the results suggest brain parenchymal entry; 5</td>
<td>[117]</td>
<td></td>
</tr>
<tr>
<td>'Migration to brain parenchyma' 'albumin modulates the transport across the blood–brain barrier'</td>
<td>Whole brain radioactivity was determined by gamma-scintigraphy of isolated mouse brain after iv. albumin-coated 15-nm ¹¹¹In-Au NP injection.</td>
<td>+Whole brain radioactivity does not verify transport across the BBB into brain parenchyma; 1.</td>
<td>21(5.1)</td>
<td>[17]</td>
</tr>
<tr>
<td></td>
<td>Brain confocal microscopic images were obtained after iv. fluorescent-labeled albumin-coated ¹¹¹In-Au NP injection</td>
<td>+Resolution is not sufficient to differentiate vascular from brain parenchymal distribution; 1</td>
<td></td>
<td></td>
</tr>
<tr>
<td>'Particles that actually crossed the BBB'</td>
<td>Gold was determined by flame AAS analysis in brain from mice that received iv. injection of 20-nm insulin PEG-coated gold particles. Brain gold was calculated from its blood concentration and brain–blood volume</td>
<td>This does not account for gold associated with BMECs that the authors hypothesize accounted for gold in the brain at late time points; 1</td>
<td>11(blood did not decrease over 48 h)</td>
<td>[118]</td>
</tr>
<tr>
<td>Coumarin distribution to the glioma region</td>
<td>Brains from glioma-bearing mice were homogenized, coumarin-6 extracted and quantified by HPLC after iv. injection of 94-nm coumarin-6, L-13RS2 ligand-coated PEG-PLGA NPs.</td>
<td>+HPLC analysis of NP component demonstrates bulk brain distribution; 1.</td>
<td>[119]</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Fluorescence spots were seen in the brain</td>
<td>+Microscopy resolution only sufficient to demonstrate brain distribution; 1</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Brain uptake</td>
<td>Radioactivity determined by gamma-scintigraphy of isolated whole mouse brain after iv. injection of 9- and 40-nm diameter, 396- and 268-nm length [¹¹¹In]-DTPA-mulitwall carbon nanotubes</td>
<td>Radioactivity demonstrates distribution of NP or component in bulk brain; 1</td>
<td>[120]</td>
<td></td>
</tr>
<tr>
<td>'Gold would presumably not be cleared from brain in such a short timeframe if it has entered the brain parenchyma'</td>
<td>Brains from mice were digested and gold determined by ICP-OES after iv. injection of 80-nm biodegradable polymeric micelles with 5 nm gold NP cores</td>
<td>ICP–OES results of NP component in bulk brain do not show brain parenchymal entry; 1</td>
<td>43(7.7)</td>
<td>[78]</td>
</tr>
</tbody>
</table>

Mean residence time from studies that reported brain and blood (in parentheses) levels at multiple times after dosing. Study multiple end points are designated by a, b, cbefore the study summary and critique. The level of demonstrated nanoparticle brain entry is from the classification described in Table 3. The numbers after the sentences in column 3 refer to the level of demonstrated NP brain entry.

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<tbody>
<tr>
<td>'Demonstrated that the NPs could cross the BBB'</td>
<td>CLSM images were obtained of brain tissue cryosections, stained with tomato lectin to label blood vessels, from rats that received iv. microbubbles and focused ultrasound to open the BBB before iv. injection of 177-nm fluorescent dye-containing PBCA NPs</td>
<td>Fluorescent spots attributed to the fluorescent dye near the blood vessels suggest NPs crossed the BBB in the presence of opened BBB; 6</td>
<td></td>
<td>[47]</td>
</tr>
<tr>
<td>'The peptide was able to diffuse outside the BBB, while NPs were restricted to the vessel endothelium'</td>
<td>CLSM images of brain sections with antibody-stained BMECs from mice iv. dosed with 35-nm fluorometric dye-labeled amphiphilic polymer-coated iron oxide NPs containing a fluorometric dye-labeled antiretroviral drug enabled distinction among microvessels, NPs and drug. The NPs and BMECs were co-localized. The antiretroviral drug was seen within and outside of the vascular compartment, in brain parenchyma</td>
<td>Peptide fluorescence was seen in brain parenchyma; 5</td>
<td></td>
<td>[24]</td>
</tr>
<tr>
<td>'Distributed into the brain tissue'</td>
<td>The capillary depletion method was used to obtain a non-BMEC fraction of mouse brains for LC-MS/MS curcin analysis after oral dosing of 139, 312 and 412 nm uncoated, chitosan- and N-trimethyl chitosan-coated, curcumin-loaded solid lipid NPs</td>
<td>Most of the whole brain curcumin was in the supernatant fraction, not associated with BMECs, but could have come from blood; 1</td>
<td></td>
<td>[71]</td>
</tr>
<tr>
<td>'BBB penetration by AuNPs'</td>
<td>Gold was determined by laser ablation ICP–MS in brain from perfused rats that received ip. injection of 1.3-nm gold NPs.</td>
<td>ICP–MS results of NP component in bulk brain do not verify BBB penetration; 2.</td>
<td>8.7(4.4)</td>
<td>[12]</td>
</tr>
<tr>
<td></td>
<td><em>Gold content was determined by ICP-MS in brain from perfused rats that received ip. injection of 1.3-nm gold NPs.</em></td>
<td>ICP-MS results of NP component in bulk brain do not verify BBB penetration; 2.</td>
<td></td>
<td>[12]</td>
</tr>
<tr>
<td></td>
<td><em>Gold was determined by laser ablation ICP-MS in brain sections</em></td>
<td>LA-ICP-MS resolution not sufficient to verify BBB penetration. 2.</td>
<td></td>
<td>[12]</td>
</tr>
<tr>
<td>'The increase of brain iron from days 1 to 7'</td>
<td>Iron was quantitated by AAS in solubilized whole organs from mice that received iv. 10-nm carboxyl-coated iron oxide magnetic NPs</td>
<td>AAS results of NP component in bulk brain do not verify BBB crossing; 1</td>
<td>Increased from 24 to 168 h after injection</td>
<td>[121]</td>
</tr>
<tr>
<td>'Exosome delivered anticancer drugs crossed the BBB and entered into the brain'</td>
<td>Fluorescent dye, paclitaxel- or doxorubicin-labeled mouse-derived &lt;50 nm BMEC exosomes were iv. injected into zebrafish transgenic for vasculature GFP expression</td>
<td>CLSM suggests NP fluorescence and shows drug fluorescence outside of the vasculature; 5</td>
<td></td>
<td>[48]</td>
</tr>
<tr>
<td>'Nose-to-brain transport'</td>
<td>Cyclosporine A was extracted from homogenized olfactory bulb and hind brain samples and quantitated by LC-MS/MS following intranasal dosing of 272-nm nanoemulsion to the rat</td>
<td>LC-MS/MS NP component shows bulk brain distribution; 1</td>
<td>2.4 and 2.6</td>
<td>[122]</td>
</tr>
<tr>
<td>'NPs were capable of penetrating cell membranes and entering cellular compartments'</td>
<td>Brains from mice with green fluorescent protein-expressing tumors were imaged using CLSM after iv. injection of 165-nm fluorescent dye-labeled poly(amineco-ester) terpolymer NPs. PECAM (in BMECs) was labeled. NPs were visualized around tumor blood vessels and within a cell</td>
<td>Brain parenchymal NP entry shown in tumor cells; 6</td>
<td></td>
<td>[49]</td>
</tr>
<tr>
<td>Brain accumulation at the site of injury</td>
<td>Whole-body IVIS images were obtained after iv. administration of approximately 170-nm fluorescent-labeled siRNA-containing biodegradable mesoporous silica, encapsulated in graphene oxide nanosheets. Fluorescence was seen in the hemisphere of mice with damaged brain</td>
<td>Greater fluorescence in brain at the injury site supports brain accumulation; 1</td>
<td></td>
<td>[123]</td>
</tr>
<tr>
<td>'NP was obviously accumulated in brain'</td>
<td>Imaging demonstrated mouse whole brain fluorescence after iv. injection of approximately 180-nm cyclic peptide-modified fluorescence dye-loaded PEG-poly(lactic acid) NPs</td>
<td>Fluorescence demonstrates brain accumulation; 1</td>
<td></td>
<td>[124]</td>
</tr>
</tbody>
</table>

Mean residence time from studies that reported brain and blood (in parentheses) levels at multiple times after dosing. Study multiple end points are designated by a, b, cbefore the study summary and critique. The level of demonstrated nanoparticle brain entry is from the classification described in Table 3. The numbers after the sentences in column 3 refer to the level of demonstrated NP brain entry.

AAS: Atomic absorption spectroscopy; BBB: Blood-brain barrier; BBTB: Blood-brain tumor barrier; BMEC: Brain microvascular endothelial cell; BSA: Bovine serum albumin; CLSM: Confocal laser scanning microscopy; EDS: Energy dispersive x-ray spectroscopy; EELS: Electron energy loss spectroscopy; FTIC: Fluorescein isothiocyanate; ICPOES: Inductively coupled plasma–optical emission spectroscopy; INAA: Instrumental neutron activation analysis; ip.: Intraperitoneal; iv.: Intravenous; IVIS: In vivo imaging system; MRT: Mean residence time; MSI: Mass spectrometry imaging; NAA: Neutron activation analysis; NP: Nanoparticle; PBCA: Poly(acrylic acid); RVG: Rabies virus glycoprotein; TEM: Transmission electron microscopy.
Table 4. Brain nanoparticle distribution study claims, summary, critique, support for the claims and level of brain or brain parenchymal entry. (cont.).

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<th>Claim</th>
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</tr>
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<tbody>
<tr>
<td>’Delivery of . . . NPs across the blood–brain barrier’</td>
<td>CLSM images of mouse brain after oral administration of 60-nm PEG-coated, curcumin and pipeline-coated glyceryl monooleate NPs showed intracellular NPs</td>
<td>Neuronal intracellular NPs supports delivery across the BBB; 6</td>
<td>18(13)</td>
<td>[50]</td>
</tr>
<tr>
<td>NP were ‘taken up into the brain cells’</td>
<td>Rats with BBB opened by focused ultrasound received iv. 22-nm fluorescent probe-labeled low-density lipoprotein NPs. Fluorescence microscopy images were interpreted as showing punctate fluorescent staining within brain cell cytosol</td>
<td>Brain parenchymal NP uptake appears to be shown in the presence of an opened BBB; 6</td>
<td>46</td>
<td></td>
</tr>
<tr>
<td>’NP retention in brain tumors’</td>
<td>*IVIS-imaged whole and ex vivo mouse brain showed accumulation in glioma after iv. 33–36-nm fluorescent-labeled, penta-amino acid- or 2-cyano-6-amino benzothiazole- or both, coated gold NP injection. *Glioma were excised for gold quantification by ICP–MS. *TEM images showed intracellular densities</td>
<td>*Whole brain results using this method show NP retention in brain tumors; 1. *ICP–MS NP component in bulk brain shows brain component retention; 1. *TEM showed brain parenchymal NP entry in tumor cells; 6</td>
<td>[125]</td>
<td></td>
</tr>
<tr>
<td>([Quetiapine fumurate]–NP (IV) may have crossed BBB by opening tight junctions)</td>
<td>Brains removed from rats were homogenized and quetiapine quantified by HPLC after iv. injection of 131-nm quetiapine fumarate-loaded chitosan NPs</td>
<td>HPLC results of NP component in bulk brain do not verify BBB crossing; 1</td>
<td>1.5(1.4)</td>
<td>[126]</td>
</tr>
<tr>
<td>Successfully overcame the in vivo BBB and BBTB delivery barriers</td>
<td>*Ex vivo brain and dissected tumors from mice with brain tumors were fluorescent imaged 4 h after iv. injection of fluorescent dye-loaded cyclic peptide-containing approximately 100-nm liposomes. *Fluorescent images of brain with fluorescent-stained microsells show fluorescence from NPs separate from microvessels</td>
<td>*Whole brain and tumor fluorescence does not demonstrate delivery across the BBB or BBTB; 1. *Spots of NP fluorescent dye distinct from blood vessels suggest NP distribution outside of brain vasculature; 5</td>
<td>[127]</td>
<td></td>
</tr>
<tr>
<td>’Ex vivo CT imaging and analysis indicated that INS-GNPs transported across the BBB’</td>
<td>*In vivo and ex vivo CT mouse brain imaging was conducted after iv. 20-nm insulin-coated, PEG-layered, gold NP administration. *Flame AAS of gold from digested brain was determined</td>
<td>*Whole brain CT images do not verify transport across the BBB; 1. *AAS results of NP component in bulk brain do not verify transport across the BBB; 1</td>
<td>15(18)</td>
<td>[128]</td>
</tr>
<tr>
<td>’Brain microvessel endothelial cell targeting’</td>
<td>Brains of mice were perfused and prepared for immunohistochemistry and fluorescence imaging after 2 iv. injections 1 h apart of 50-nm transferrin antibody-conjugated mesoporous silica NPs. BMECs were labeled with biotinylated potato lectin detected by fluorescent-labeled streptavidin. One hour after the second injection NPs were associated with the BMECs. There was no evidence brain parenchymal penetration</td>
<td>NP brain microvessel targeting was demonstrated; 3</td>
<td>[129]</td>
<td></td>
</tr>
<tr>
<td>’Enhanced drug transport across BBB’</td>
<td>*Iron was determined by AAS in mouse organs after iv. injection of 80-nm dextran-spermine superparamagnetic-core NPs and 96-nm transferrin-coated NPs *Tissues were stained with Prussian blue to visualize iron localization</td>
<td>*AAS results of NP component in bulk brain do not verify transport across the BBB; 1</td>
<td>Increased from h 24 to 168 after injection</td>
<td>[130]</td>
</tr>
<tr>
<td>’To further confirm the translocation of CLS-PEG NPs across the BBB’</td>
<td>*Rat brains were analyzed by fluorescence analysis after iv. injection of 176-nm coumarin-6-labeled cholesterol-containing PBCA NPs and 185-nm particles also containing PEG. *Brains from perfused rats were viewed using CLSM that showed uniform extracellular fluorescence</td>
<td>*Fluorescence results do not confirm BBB translocation; 1. *NP component in brain extracellular space supports claim of its passage through the BBB; 5</td>
<td>2.2 and 2.3(3.4 and 5.5)</td>
<td>[52]</td>
</tr>
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Mean residence time from studies that reported brain and blood (in parentheses) levels at multiple times after dosing. Study multiple end points are designated by a, b, c before the study summary and critique. The level of demonstrated nanoparticle brain entry is from the classification described in Table 3. The numbers after the sentences in column 3 refer to the level of demonstrated NP brain entry.
Table 4. Brain nanoparticle distribution study claims, summary, critique, support for the claims and level of brain or brain parenchymal entry. (cont.).

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</tr>
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<tbody>
<tr>
<td>'The NPs crossed the BBB and entered the brain parenchyma'</td>
<td><em>Gold in brains from perfused rats was determined by INAA after iv. injection of rhodamine-PEG-angiopep-2-labeled gold nanorods. B Brain from a rat was fluorescent imaged after iv. injection of rhodamine-PEG-angiopep-2-labeled gold nanorods. TEM of hippocampus of a perfused rat after iv. injection of PEG-angiopep-2-labeled gold nanorods demonstrated intracellular electron dense spots.</em></td>
<td>INAA results of NP component in bulk brain does not demonstrate distribution across the BBB; 1. Fluorescence results do not confirm BBB translocation; 1. NP seen in brain parenchyma; 6. Assuming the spots were gold nanorods</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Brain delivery

| Brain delivery | Brains from mice were homogenized for LC/MS RU determination after iv. injection of marigold flower extract (RU)-, polysorbate-80-containing lipid polymer hybrid PLGA NPs | LC/MS results of NP component support bulk brain delivery; 1 | 2.5(1.2) | [132] |

Uptake by neuronal cells and appreciable crossing of blood-brain barrier

| Uptake by neuronal cells and appreciable crossing of blood-brain barrier | Brains from rats were homogenized, solvent extracted and dimethyl fumarate determined by HPLC after oral administration of dimethyl fumarate-containing solid lipid NPs | HPLC results of NP component in bulk brain do not verify neuronal cell uptake or BBB crossing; 1 | 5.7(4.7) | [133] |

Lamotrigine brain uptake

| Lamotrigine brain uptake | Lamotrigine was determined by HPLC in homogenized brain after iv. injection of 125-nm lamotrigine-loaded poly-ε-(D,L-lactide-co-caprolactone) NPs | HPLC results of NP component show bulk brain uptake; 1 |  |  |

[DNA-coated gold NPs] can be successfully delivered to specific brain locations across the BBB in mice using focused ultrasound

| [DNA-coated gold NPs] can be successfully delivered to specific brain locations across the BBB in mice using focused ultrasound | Mice were iv. injected with 6-14-nm fluorescent-labeled DNA-coated gold NPs after unilateral microbubble injection and before ultrasound to open the BBB. The effected hemisphere showed more NPs (bright-field and fluorescence microscopy) than the contralateral hemisphere | Resolution was not sufficient to verify delivery across the BBB; 1 |  |  |

This strategy increased NP translocation into brain parenchyma

| This strategy increased NP translocation into brain parenchyma | CLSM of brain, stained for GFAP to localize astrocyte foot processes, from mice iv. injected with 180-nm ApoE4-coated, fluorescent dye-labeled, lipid, polysorbate-80 NPs showed some fluorescence from the NPs not surrounded by fluorescence from GFAP | Fluorescence from the NP label appears to be in brain parenchyma; 5 |  |  |

Gamma scintigraphic images showed ... the radiolabeled formulations successfully crossed BBB

| Gamma scintigraphic images showed ... the radiolabeled formulations successfully crossed BBB | *Whole rat gamma scintigraphy was conducted after iv. injection of 24-nm AZT-containing 99mTc-labeled nanoliposomes. B AZT was determined by HPLC in homogenized brain of in situ blood-perfused rats after iv. injection of 24 nm AZT-containing 99mTc-labeled nanoliposomes | *Whole rat radioactivity does not verify brain entry or BBB crossing. HPLC results of NP component in bulk brain do not verify BBB crossing 2 | 6.2(8.8) | [136] |

Blood-brain barrier permeability of the NPs was successfully demonstrated by the incorporation of fluorescent dye, Rhodamine B in the NPs

| Blood-brain barrier permeability of the NPs was successfully demonstrated by the incorporation of fluorescent dye, Rhodamine B in the NPs | Brain from rats was homogenized, solvent extracted and Rhodamine B determined by HPLC after ip. injection of 300–400-nm saxagliptin-loaded chitosan-L-valine-based NPs | HPLC results of NP component in bulk brain do not verify BBB permeability; 1 |  |  |

Brain accumulation of curcumin-loaded NPs

| Brain accumulation of curcumin-loaded NPs | Fluorescence in homogenized brains from perfused rats was determined by HPLC after iv. injection of 46-nm 6-coumarin-loaded oleic acid-conjugated PEG-poly-(epsilon-caprolactone) NPs | HPLC results of NP component show bulk brain delivery; 2 |  |  |

After intravenous administration, fluorescent [solid lipid NPs] were detected in brain parenchyma outside the vascular bed, confirming their ability to overcome the BBB

| After intravenous administration, fluorescent [solid lipid NPs] were detected in brain parenchyma outside the vascular bed, confirming their ability to overcome the BBB | Epi-fluorescence microscopy of rat brain 24 h after iv. injection of 278-nm FITC and Brij 78-containing solid lipid NPs showed spherical fluorescence not associated with an endothelial marker | Spherical fluorescence in absence of blood vessel marker fluorescence suggests NP brain parenchymal entry; 5 |  |  |

Mean residence time from studies that reported brain and blood (in parentheses) levels at multiple times after dosing. Study multiple end points are designated by *h < before the study summary and critique. The level of demonstrated nanoparticle brain entry is from the classification described in Table 3. The numbers after the sentences in column 3 refer to the level of demonstrated NP brain entry.

Ex vivo NP delivery to the brain via the nose-to-brain pathway demonstrated delivery of drugs to the brain via the nose-to-brain pathway 'rapidly and effectively delivered drugs to the brain via the nose-to-brain pathway'.

Bioavailability of quercetin in the brain

Brain distribution/delivery

Penetration of NPs into brain parenchyma

NPs 'passed the BBB, and entered the cytoplasm of vascular endothelial cells, as depicted by the biodistribution studies, flow cytometry analysis and subcellular TEM micrographs.'

'NP formulation increased BBB transport ... in vivo'

Enhanced delivery of [doxorubicin] across the BBB

'NP' rapidly and effectively delivered drugs to the brain via the nose-to-brain pathway

NP delivery to the brain

Table 4. Brain nanoparticle distribution study claims, summary, critique, support for the claims and level of brain or brain parenchymal entry. (cont.)

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<tbody>
<tr>
<td>'Ex vivo imaging and semiquantitative assay of the brain parenchymal entry' NPs' penetration behavior of ... NPs'</td>
<td>Whole mice were multispectral and ex vivo brain CLSM imaged after iv. injection of 148-nm menthol BSA NPs. CLSM and TEM images were obtained of the pineal gland (which lacks a BBB) of rats after iv. injection of 148-nm menthol BSA silver NP injection</td>
<td>Whole-body multispectral and CLSM imaging show bulk brain delivery; 1</td>
<td>[53]</td>
<td></td>
</tr>
<tr>
<td>Bioavailability of quercetin in the brain</td>
<td>Brains from rats were homogenized, proteins precipitated and quercetin determined by HPLC after oral feeding of 30–50-nm quercetin-coated superparamagnetic iron oxide NPs</td>
<td>HPLC results of NP component show bulk brain delivery; 1</td>
<td>[140]</td>
<td></td>
</tr>
<tr>
<td>Brain distribution/delivery</td>
<td>Brains from rats were homogenized, proteins precipitated and rivastigmine determined by HPLC after iv. injection of 145-nm rivastigmine-loaded mesoporous silica NPs</td>
<td>HPLC results of NP component show bulk brain delivery; 1</td>
<td>6.5</td>
<td>[141]</td>
</tr>
<tr>
<td>Penetration of NPs into brain parenchyma</td>
<td>Three adjacent sections from isolated guinea pig brains were assessed by silver-enhanced brightfield microscopy, CLSM and TEM after arterial perfusion for 120 min of approximately 8 nm fluorescent-labeled mercaptopoundecysulfonic acid/octanethiol coated gold/iron core NPs followed by 30-min washout perfusion</td>
<td>CLSM indicates fluorescence surrounding cell nuclei. Optical microscopy shows silver stain surrounding blood vessels. TEM shows electron dense particles on the luminal wall of BMECs and inside brain cells; 6</td>
<td>[22]</td>
<td></td>
</tr>
<tr>
<td>NPs 'passed the BBB, and entered the cytoplasm of vascular endothelial cells, as depicted by the biodistribution studies, flow cytometry analysis and subcellular TEM micrographs.'</td>
<td>Brains from perfused mice were homogenized and cells analyzed by flow cytometry 1 h after ip. injection of fluorescent-labeled glucose- or PEG-glucose-coated silica NPs. Brains from perfused mice were CLSM imaged 1 h after ip. injection of fluorescent-labeled glucose- or PEG-glucose-coated silica NPs.Brains from harvested mice were TEM imaged 1 h after ip. injection of fluorescent-labeled PEG-glucose-coated silica NPs.</td>
<td>Flow cytometry does not reveal cell type or NP localization in, vs on, the cell; 2CLSM images do not verify brain parenchymal localization; 2TEM shows electron densities, perhaps in neurons; 6</td>
<td>[54]</td>
<td></td>
</tr>
<tr>
<td>'NP formulation increased BBB transport ... in vivo'</td>
<td>Whole mouse fluorescence imaging was conducted from 0.25 to 24 h after intranasal dosing of indoxyamine green containing transferrin or RVG-coated BSA NPs. Oxytocin in CSF was determined by enzyme-linked immunosorbent assay after intranasal dosing of oxytocin-containing RVG-coated BSA NPs.</td>
<td>Whole body bioimaging does not verify BBB transport; 1NP component in CSF does not demonstrate brain uptake</td>
<td>7.4</td>
<td>[82]</td>
</tr>
<tr>
<td>'Enhanced delivery of [doxorubicin] across the BBB'</td>
<td>HPLC of dialysates from microdialysis of rat hippocampus after intranasal dosing of 92-nm polysaccharide NP containing doxorubicin showed doxorubicin concentration peak at 30 min</td>
<td>Peak doxorubicin concentration this soon cannot be due to nose to hippocampus neuronal transport. It would be distribution across the BBB. NP component does not demonstrate brain NP uptake; 1</td>
<td>4.2</td>
<td>[77]</td>
</tr>
<tr>
<td>'NP' rapidly and effectively delivered drugs to the brain via the nose-to-brain pathway'</td>
<td>Mouse brain was IVIS imaged ex vivo and MRI imaged after intranasal dosing of 110- or 143-nm fluorescent and gadolinium-labeled cholera toxin or BSA NP</td>
<td>Whole brain results using these methods show fluorescence and gadolinium in the brain; 1</td>
<td>11.6</td>
<td>[142]</td>
</tr>
<tr>
<td>NP delivery to the brain</td>
<td>Ex vivo brain was IVIS imaged after iv. injection of fluorescent-labeled, microbubble complexed, albumin NPs and ultrasound to brain tumor bearing mice</td>
<td>Whole brain results using this method show fluorescence in the brain; 1</td>
<td></td>
<td>[143]</td>
</tr>
</tbody>
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Mean residence time from studies that reported brain and blood (in parentheses) levels at multiple times after dosing. Study multiple end points are designated by a,b,c before the study summary and critique. The level of demonstrated nanoparticle brain entry is from the classification described in Table 3. The numbers after the sentences in column 3 refer to the level of demonstrated NP brain entry.

AAAS: Atomic absorption spectroscopy; BBB: Blood-brain barrier; BBTB: Blood-brain tumor barrier; BMEC: Brain microvascular endothelial cell; BSA: Bovine serum albumin; CLSM: Confocal laser scanning microscopy; EDS: Energy dispersive x-ray spectroscopy; EELS: Electron energy loss spectroscopy; RTIC: Fluorescein isothiocyanate; ICP-MS: Inductively coupled plasma–mass spectrometry; ICP-OES: Inductively coupled plasma–optical emission spectroscopy; INAA: Instrumental neutron activation analysis; ip.: Intraperitoneal; iv.: Intravenous; IVIS: In vivo imaging system; MRT: Mean residence time; MSI: Mass spectrometry imaging; NAA: Neutron activation analysis; NP: Nanoparticle; PBCA: Poly(n-butyl cyanoacrylate); RVG: Rabies virus glycoprotein; TEM: Transmission electron microscopy.
dependent, increasing from less than 96 h after 60 μg to approximately 480 h after 1600 μg [81], suggesting the quantity of NPs in brain parenchyma influence their duration of persistence in that compartment.

Some studies reported the brain level of the NP or its components at multiple times after its peripheral administration. Some studies also reported blood and/or peripheral organ levels. The reported values are shown in Supplementary Figure 1. To be able to visually compare the rates of NP/component increase or decrease over time given the wide range of levels and timecourses among the studies, the results were normalized to the dose for CED studies or first reported value for other studies and shown as double log figures. It is expected that NPs that entered brain parenchyma would persist longer in the brain than NPs that did not cross the BBB, and longer than in the blood. Most reports of brain NP or component determination over time did not have many values (range for the non-CED and non-microdialysis studies: 2–16, median: 5) discouraging rigorous pharmacokinetic analysis or estimate of NP residence to infinity. Noncompartmental mean residence time (MRT) to the last sample was determined using Phoenix 8.1 WinNonLin. The MRT of NPs or their components after peripheral or intranasal administration is generally less than after CED, suggesting less than 100% of the NPs or components entered brain parenchyma. Good agreement was obtained between the MRT results and author reported half-lives for the same datasets (Tables 1 & 4). The MRT was similarly calculated for NP, or its component, in blood from the studies that determined them at multiple times after NP administration. The results are shown in parentheses in Table 4. The timecourse and MRTs for the NP or its component in the brain, blood and peripheral organs were often similar (Supplementary Figure 1 & Table 4). This could be due to most or all of the NP/component in the brain residing in blood within the brain or adhering to the luminal wall of BMECs and being cleared from brain at the same rate as blood clearance from peripheral circulation. Exceptions were seen in two reports where the brain MRT was considerably longer than blood MRT [17,58], suggesting NP distribution beyond the vascular compartment. However, evidence of brain parenchymal entry was not provided.

Studies classified according to the criteria in Table 3 as demonstrating brain parenchymal entry (levels 5 and 6) that reported brain NP or component levels at multiple times are shown in Supplementary Figure 1D and E. Brain levels often increased over time or decreased more slowly than in studies classified as not showing brain parenchymal entry. Brain levels often decreased more slowly than blood levels in studies shown in Supplementary Figure 1D

Table 4. Brain nanoparticle distribution study claims, summary, critique, support for the claims and level of brain or brain parenchymal entry. (cont.).

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</tr>
</thead>
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<tr>
<td>NP ‘transport across the BBB after intravenous administration’</td>
<td>CLSM images are claimed to show 140–200-nm phosphatidic and sialic acid, 5-hydroxytryptamine-modulin-modified PLGA NPs in the hippocampus of rats after their iv. injection</td>
<td>No fluorescence attributed to the NPs can be seen</td>
<td>[55]</td>
<td></td>
</tr>
<tr>
<td>BBB penetration and NP distribution into brain parenchyma</td>
<td>Epifluorescence was determined in brain regions that did not contain vasculature after iv. injection of 4-nm Angiopep-2, transferrin receptor ligand, endosome escape unit, fluorescent marker containing malic acid NPs in mice whose cerebrovascular was stained. The results were interpreted to indicate that the NPs traversed the BBB and approximately 40% of plasma fluorescence was in the parenchyma with the highest concentration proximal to the vasculature</td>
<td>The methods suggest the fluorescent component of the NP entered brain parenchyma; 5</td>
<td>[144]</td>
<td></td>
</tr>
<tr>
<td>[NP] crossing intact BBB was ... demonstrated in healthy mice’</td>
<td>Confocal microscopy of mouse brain after iv. injection of 126-nm doxorubicin, mitomycin C and internalizing peptide-labeled terpolymer-lipid hybrid NP after iv. injection of Texas-red-dextran to visualize blood vessels showed particulate doxorubicin fluorescence not associated with blood vessels</td>
<td>Punctate fluorescence attributed to doxorubicin not associated with blood vessels suggests brain parenchymal NP entry; 6</td>
<td>[145]</td>
<td></td>
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Mean residence time from studies that reported brain and blood (in parentheses) levels at multiple times after dosing. Study multiple end points are designated by a, b, c before the study summary and critique. The level of demonstrated brain nanoparticle entry is from the classification described in Table 3. The numbers after the sentences in column 3 refer to the level of demonstrated NP brain entry.

and E than B or C. There is some concordance between the classification levels and NP/component timecourse in the brain, suggesting the rate of NP/component clearance from the brain reflects the absence or presence of brain parenchymal entry.

The evidence of NP parenchymal entry may not be from the same measure as that providing the brain timecourse (e.g., TEM showing NP in brain cells, whereas NP/marker residence determined by HPLC). When NP or component was determined by radioactivity, neutron activation analysis, IVIS imaging, fluorescence, HPLC, atomic absorption spectroscopy, inductively coupled plasma–optical emission spectroscopy, inductively coupled plasma–mass spectrometry or LC–MS, it could be that only a fraction was in brain parenchyma. A risk of concluding NP brain entry or brain parenchymal entry from a component of the NP, such as an incorporated fluorescent dye or a metal component of a metal- or metal oxide-containing NP, is that the dye or metal might be released and its distribution does not represent the NP. For example, it was concluded that dye clearance from the brain in 24 h was ‘most likely due to degradation of the dye itself rather than actual clearance of the NPs from the brain’ [82]. Similarly, metal analysis by neutron activation analysis, atomic absorption spectroscopy, inductively coupled plasma–optical emission spectroscopy or inductively coupled plasma–mass spectrometry may be informing about released ion, particularly from readily soluble NPs such as those containing copper, silver and zinc.

In contrast, the results of many studies which did not demonstrate NP distribution in brain parenchyma showed more rapid clearance of brain-associated NPs (Table 4 & Supplementary Figure 1). Many studies did not remove blood from the brain. The decline of NPs within a few hours in the brain, at a rate similar to its decline in blood, and more rapidly than the decline from peripheral organs, may be due to lack of brain parenchymal entry. Such results suggest that only some, or none, of the NPs crossed the BBB. The NPs may have adhered to the luminal wall of brain vasculature or localized in cellular or membrane components of the BBB, and subsequently distributed away from these sites into blood circulating through the brain. In studies that did not use sufficient methods to demonstrate NP brain parenchymal entry, the NPs may indeed have crossed the BBB. However, the methods do not verify that. If procedures are used to rule out the contribution of NPs in blood and adherence to BMECs, NP persistence in the brain may indicate successful distribution across the BBB.

**Conclusion**

The BBB and BCSFB provide a significant challenge to deliver substances into the brain to access neural, glial and cancer cells. Verification that the substance has entered into the compartment that houses these cells (brain parenchyma) is not trivial. Experimental results are sometimes overinterpreted to suggest successful delivery of substances that might directly act on cells in the brain when verification of brain parenchymal entry is lacking. An understanding of the anatomy and location of the barriers (BBB and BCSFB) to brain parenchymal entry helps one interpret if a substance has entered brain parenchyma. Visual and procedural methods have been used to verify brain parenchymal entry. In this article, methods have been classified as to the level of verification of brain parenchymal entry and then critically applied to studies. The BBB can be bypassed by direct substance introduction into the brain by CED that often results in prolonged residence of the substance in the brain. It is suggested that prolonged substance residence in the brain after its peripheral (e.g., iv.) administration suggests brain parenchymal entry. It is hoped that this guide will help researchers and reviewers be more precise in their demonstration and interpretation of substance distribution in the brain.

**Future perspective**

As the study of NP biological interaction matures, there is greater understanding of how to critically assess NP effects and their distribution. If the criteria and methods that demonstrate bulk brain and brain parenchymal entry proposed herein are employed by researchers and reviewers, there will be more precise reporting of brain NP distribution. More realistic result interpretation will enable more realistic clinical benefit expectation. This will enhance the credibility of this endeavor and help investigators engage in best practice research.

**Supplementary data**

To view the supplementary data that accompany this paper please visit the journal website at: www.futuremedicine.com/doi/suppl/10.2217/nnm-2019-0169
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No writing assistance was utilized in the production of this manuscript.

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Executive summary

- The anatomy and function of the blood–brain (BBB) and blood-cerebrospinal fluid barriers are reviewed.
- Exposure routes to bypass the BBB (convection-enhanced delivery and uptake from the nasal cavity by cranial nerves) are described.
- Visual and procedural methods to determine nanoparticle (NP) brain and brain parenchymal entry are presented.
- Criteria are proposed to classify levels of experimental result evidence that demonstrate NP bulk brain or brain parenchymal entry.
- Studies of NP brain entry or distribution across the BBB are summarized, critiqued and evaluated using the classification levels to interpret if the study shows NP bulk brain or brain parenchymal entry.
- NP or component persistence in the brain after systemic or intranasal administration compared with blood and peripheral organs and direct infusion by convection-enhanced delivery into the brain are compared with interpret support for NP distribution beyond the brain vasculature, into brain parenchyma.

References

Papers of special note have been highlighted as: ● of interest; ●● of considerable interest


Demonstration that nanoparticles (NPs) in blood within the brain can have a considerable contribution to bulk brain NPs.


• A perfusion method to remove blood from the brain.


• An electron micrograph figure showing NPs adhering to the luminal wall of brain microvascular endothelial cells after perfusion to remove blood from the brain.


• One of the best demonstrations differentiating NP and NP component distribution within and not within brain parenchyma.


• One of the best demonstrations differentiating NP and NP component distribution within and not within brain parenchyma.


**One of the best demonstrations differentiating NP and NP component distribution within and not within brain parenchyma.**


- The timecourse of NP component in the brain compared with blood suggests distribution out of blood, perhaps into brain parenchyma. Imaging results support this suggestion.


