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**Blood-brain barrier regulation**

**Environmental cues controlling the onset of barrier properties**

Mark Ronald Mizee and Helga Eveline de Vries*

Department of Molecular Cell Biology and Immunology (MCBI); Neuroscience Campus Amsterdam; VU University Medical Center; Amsterdam, The Netherlands

The existence of a barrier between the central nervous system (CNS) and the systemic circulation has been described over one hundred years ago. Since the discovery that this barrier was the result of specific barrier properties of the brain endothelial cells, research has focused on the identification of pathways that instruct brain endothelial cells to form the highly specialized blood-brain barrier (BBB). Even though our current understanding of BBB development is far from complete, recent literature shows a rise in knowledge of CNS-specific cues that can drive BBB development.

In this commentary, we will provide a brief overview of brain selective factors that are critical in the development of barrier properties in the brain endothelium; in particular the role of retinoic acid will be discussed.

**The Blood-Brain Barrier**

The vasculature of the brain functions as a specialized barrier to protect the central nervous system (CNS) from the systemic circulation by restricting entry of unwanted molecules and immune cells into the brain, and by active removal of cytotoxic compounds. Importantly, the endothelial cells that line the lumen of the cerebral vessels are also equipped to actively supply the brain with essential nutrients and oxygen through specific transport mechanisms. The blood-brain barrier (BBB) is not a rigid barrier but a dynamic structure that receives continuous input from the CNS cells it protects. This allows for a thorough response to the local demands for oxygen, nutrients, and buffering which is crucial for the maintenance of a CNS homeostasis that favors optimal neuronal function.1-4

The blood-brain barrier (BBB) is composed of highly specialized brain endothelial cells (ECs) that line the vascular wall of the brain capillaries. Brain ECs form a tight structural barrier by the polarized assembly of tight junction (TJ) proteins at the luminal side of the endothelium and a metabolic barrier by the expression of membrane efflux pumps. Brain ECs are enclosed together with pericytes within the basement membrane onto which astrocytes firmly project their endfeet. Combined with the input of neurons and microglia this structure forms the neurovascular unit (NVU), which ensures optimal protection of the CNS from harmful compounds and the close regulation of CNS homeostasis.1-4

A schematic overview of the NVU in the healthy CNS is depicted in Figure 1.

The BBB limits both transcellular and paracellular passage of cells and molecules from the systemic circulation into the CNS and vice versa. Transcellular passage of hydrophilic molecules is limited due to a low rate of transcytotic vesicles, an extremely low pinocytotic activity, expression of active efflux membrane pumps of the ATP-binding cassette (ABC) family such as P-glycoprotein, and high metabolic activity (cytosolic enzymes and transporters).5-7 To buffer excess amounts of neurotransmitters like glutamate from the CNS, brain ECs possess excitatory amino acid transporters (EAAT) 1–3, to limit neurotoxicity. In order to closely regulate the influx of only those components that are necessary...
in the CNS, brain ECs harbor specific transporters that actively transport nutrients like glucose into the CNS by glucose transporters (GLUT1–3).14

Paracellular diffusion of hydrophilic molecules and trafficking of immune cells is restricted by a network of TJ complexes which allow firm adhesion of brain ECs to each other and sealing of the interendothelial space. Adjacent brain ECs express continuous rows of transmembrane proteins that make homophilic contact in the intercellular space and form TJs. Claudins and occludin are the most important transmembrane components of TJs, but the participation of junctional adhesion molecules (JAMs) and adherens junctions (Cadherins) are important as well. The C-terminal cytoplasmic domain of occludin, the first described TJ protein, is associated with the zona occludens (ZO)-1 and ZO-2 proteins, which link occludin to the cytoskeleton. Claudins make up a family of proteins that consists of at least 23 closely related members. At the BBB, the presence of claudin-1,-3,-, 5 and recently -12 has been reported.8-11

The endothelium of the CNS microvasculature shows a high degree of specialization to form the BBB, and the regulatory process behind this specialization is still largely unknown. However, in the past decade, a number of pathways involved in the development and maintenance of the brain ECs have been put forward.

Cells Involved in BBB Functioning

CNS cells surrounding the endothelial layer are thought to provide angiogenic ECs with the appropriate signals for BBB maturation, as well as with the signals required for maintenance of the mature BBB-phenotype. The different cell types involved in the regulation of the endothelial cell phenotype as observed in the BBB are described below.

Astrocytes

Astrocytes are strongly represented within the neurovascular unit, ensheathing over 95% of the abluminal microvascular surface. It was this observation that gave rise to the idea that astrocytic processes formed the BBB, until electron microscopic studies showed that brain ECs were responsible for barrier function in brain microvasculature.

Astrocytes are able to influence a number of features of the brain ECs, leading to increased integrity of the BBB. TJ expression and TJ complex formation and maturation, expression and localization of brain EC transporters, and specialized enzyme systems have been shown to be upregulated under astrocyte influence. The notion that astrocytes can induce and maintain BBB properties in brain ECs through physical interaction and secreted agents has been widely accepted. Astrocyte processes extending toward CNS microvessels terminate in specialized (perivascular) endfoot structures onto the basal lamina surrounding the brain ECs. Astrocyte endfeet associated with brain ECs show a high density of orthogonal arrays of particles (OAPs), organized arrays of ion- and volume-regulating membrane particles identified by freeze fracture, containing channels like the water channel aquaporin-4 (AQP4) and the potassium ion channel Kir 4.1.12 Membrane proteins in OAPs represent a strong polarization of perivascular astrocyte function and correlate with the expression of the basement membrane molecule agrin, an important proteoglycan for BBB integrity,13 responsible for the correct localization of AQP4 (see Fig. 1). The distribution of these channels in OAPs is most likely important in the regulation of BBB homeostasis, as disruption of this distribution is associated with microvascular damage in neuropathologies such as Alzheimers disease.14

The observation of astrocyte conditioned medium inducing junction formation in brain ECs in vitro gave rise to the idea that astrocyte-derived secreted factors were able to influence the BBB properties of brain ECs. Numerous astrocyte-derived agents have since then been described, mainly by in vitro studies, as modulators of brain EC barrier function. Transforming growth factor-β (TGFβ) secreted by astrocytes has been shown to mediate the regulation of tissue plasminogen activator and the anticoagulant thrombomodulin.17 Glial-derived neurotrophic factor (GDNF) has been found to enhance barrier function in brain ECs through the induction of TJ expression.18 Fibroblast growth factor (FGF) was found to decrease BBB permeability9 consistent with the observation that FGF knockout mice show decreased levels of TJ proteins and BBB integrity loss. Furthermore, FGF was shown to be an astrocyte secreted factor that increases TJ expression of brain ECs with an important effect on BBB permeability.19,20 Recently, sonic hedgehog (Shh), a member of the Hh pathway, was shown to be produced and secreted by perivascular astrocytes in the human and mouse adult brain and that microvascular brain ECs expressed the receptors and the intracellular machinery to respond to Hh ligands. Pharmacological neutralization of Hh receptors or genetic deletion of Hh receptors lead to enhanced permeability of the BBB and loosening of the TJs.21 Together, these observations confirm the important role of perivascular astrocytes in the regulation of the BBB in the adult CNS.

Pericytes

Pericytes are perivascular, contractile cells that closely associate with capillary walls, and directly contact the brain EC membrane.22 Pericytes are thought to exert influences on the brain EC, through their specialized junctions, involving gap junctions, TJs, and AJs.23,24 Although the molecular mechanism by which pericytes mediate vascular integrity is not yet understood, perivascular pericytes are known to release growth factors and angiogenic molecules which are able to regulate microvascular permeability and angiogenesis. Besides influencing brain EC function, pericytes also contribute to the stability of microvessels and cover a large part of the abluminal brain EC surface, further influencing BBB permeability.23,24 The regulation of blood flow in CNS capillaries by pericytes has been shown to result from pericytes contracting and relaxing in a regulated manner.

Neurons

Due to the high metabolic need of neurons and the dynamic pattern of neural activity, the CNS requires a tight regulation of the microcirculation which provides the necessary nutrients and means of waste transport. The coupling of brain activity and CNS blood flow is
therefore crucial for normal neuronal functioning. Although the cellular aspect of this coupling is not fully understood, the involvement of all components of the neurovascular unit seems to be necessary for the regulation of CNS blood flow by neurons. Besides the indirect regulation of blood flow, neurons are also found to directly innervate brain EC or brain EC-associated astrocytes functioning as a liaison for neuronal-endothelial coupling. Because disruption of BBB integrity is often found to accompany pathological changes in CNS blood flow, it was suggested that the observed BBB permeability changes were due to active involvement of neurons in BBB integrity. Indeed, noradrenergic, serotoninergic, cholinergic, and GABA-ergic neurons have been found to directly contact the microvascular endothelium. Although the mechanism of action is unknown, neurons innervating the neurovascular unit are thought to regulate BBB permeability. An example of this regulation is shown by the loss of cholinergic innervation of the CNS microvasculature, resulting in impaired cerebrovascular functioning in AD. In short, neurons in the NVU do not only play an active part in the regulation of CNS blood flow, but also seem able to directly influence BBB permeability, through direct innervations of brain EC.

**Development of the Blood-Brain Barrier**

The vascular system of the CNS arises early in embryogenesis through the invasion of vascular plexus-forming angioblasts into the head region, followed by invasion of the CNS by vascular sprouts from the peri-neural vascular plexus, extending toward the ventricles. Peripheral vascular system development has been described in detail and various signaling systems taking part in vasculogenesis, angiogenesis, and differentiation have been uncovered. However, few reports exist on developmental CNS-specific cues for the induction of the specialized EC phenotype found at the BBB.

**Pericytes**

Pericytes have recently emerged as a major contributor to the development of the BBB during embryogenesis, showing increased permeability of the BBB and dysregulation of junction protein localization in the microvasculature of pericyte-deficient mice. Although
contradictory evidence exists regarding
the induction of brain EC-specific
gene expression patterns by pericytes,
the presence of these perivascular cells,
thought to originate from neural crest
cells, is crucial in the very early stage
of BBB development. Since many
developmental events occur in parallel at
the immature BBB, it seems likely that
the interplay of radial glia, pericytes,
and neural progenitors is needed for
the correct patterning and subsequent
maturation of the BBB. Interestingly,
pericyte-deficiency during development
also leads to mislocalization of astrocyte-
endfeet on the endothelial basal lamina
and a disturbed polarization of endfeet-
specific proteins.32,33

**Radial glia**

During CNS development, radial glial
cells provide structural and trophic cues
and in the later stages of development
differentiated astrocyte endfeet projections
provide an almost complete enveloping
of the brain microvasculature in adult vertebrates.34 The search for CNS-specific signals which affect the BBB phenotype in brain ECs has implicated astrocytes and glial progenitors as inducers of a
specific BBB-phenotype in brain EC.
Interestingly, the developmental window
in which the vasculature invades the
developing CNS and matures into the
BBB overlaps with the induction of
neuronal differentiation and outgrowth
and both systems share guidance cues
and differentiation-inducing signaling
pathways.35 Radial glial cells have a
prominent role in neuronal differentiation
and outgrowth, leading to the hypothesis
that these cells also function as BBB-
inducing cells during development. This
is illustrated by the fact that until recently,
sonic hedgehog (Shh)-signaling in the
CNS has been mostly associated with
neuronal patterning and differentiation.
However, Shh was shown to be released by
fetal astrocytes, and able to induce BBB-
properties in ECs during embryonic CNS
development.21

**Wnt-signaling**

CNS-specific Wnt/β-catenin signaling
has been firmly implicated in normal
BBB-development. Animal models in
which β-catenin activation was ablated
showed decreased BBB-maturation
and increased permeability, whereas no
effects were reported on the non-CNS
vasculature. Furthermore, the expression
of Wnt-ligands associated with canonical
β-catenin activation in ECs where shown
to be region dependent and only detectable
in neural progenitors.36-38

**Morphogens and BBB onset: role for
retinoic acid**

The recent finding that a morphogen
like Shh induces BBB properties, paved
the way for the investigation of other well-
known neuronal differentiation signals, in
view of BBB development. Considering
the overlap in guidance and differentiation
cues of both the neuronal and vascular
systems during CNS development, and
the involvement of radial glial cells in
both processes, we hypothesized that the
morphogen retinoic acid (RA) may have
a role in BBB onset. Retinoic acid (RA) is
a powerful vitamin A-derived morphogen
in early CNS development and radial
glia-derived RA is crucial in normal
neurogenesis.

Upon our recent publication,39 the
involvement of RA in BBB development
during CNS embryogenesis had not been
investigated. Using our in vitro models for
the human BBB, we first described that
astrocyte-conditioned medium (ACM)
derived from fetal astrocyte cultures affects barrier properties. We furthermore showed that the enzyme responsible for RA-production, retinaldehyde dehydrogenase (RALDH), is expressed by radial glial cells in the developing human CNS. The release of RA by radial glial cells has also been shown by others, as well as the dependence of neural progenitors on radial glia-derived RA as a differentiation factor. However, our group was the first to report on the effect of radial glia-derived RA on brain ECs and BBB development, which is summarized in Figure 2.

Combined with the recent report of sonic hedgehog (Shh)-induced barrier formation,21 these findings reflect the importance of astrocyte precursors in the induction of the BBB during CNS development. Another important consideration is the difference between human and mouse studies, when investigating developmental mechanisms. Most developmental studies incorporate the detailed investigation of the murine CNS, which does not always extrapolate well to the human situation, as exemplified by species-differences in CNS drug discovery. We therefore also investigated the developing human CNS, in which microvascular responsiveness to RA was shown by the presence of endothelial RA receptor-β during the onset of barrier formation. Even though this is not final proof, it does indirectly show the involvement of RA signaling of BBB induction in the human CNS.

Further investigation of the effects of RA on human brain ECs in vitro showed a positive regulation of BBB-characteristics like high electrical barrier resistance combined with an increased expression of genes associated with high paracellular and transcellular resistance. However, in vitro studies using brain ECs and varying culture conditions like cell-conditioned media, astrocyte and pericyte co-culture, or endogenous and exogenous ligands have led to a plethora of soluble molecules that are reported to increase brain EC-function or barrier-related gene expression. Whether all described soluble factors are involved in either development or maintenance of the BBB however, cannot be based on in vitro studies alone.39 We therefore investigated the effect of RA receptor inactivation on BBB formation during a specific time window (E10-E16) in the developing murine brain. The most important finding in our in vivo experiments was the profound leakage of immunoglobulins and fluorescent tracer into the brain parenchyma of treated mice. Since inhibiting the RA pathway might also affect other described BBB inducing signals, the interplay of the identified pathways in the distinct time frames during embryogenesis and the downstream effects on the BBB development need to be elucidated.

**Perspective**

A better understanding of all developmental systems converging on brain ECs during BBB development does not merely answer fundamental scientific questions. New insights in BBB development might also provide new avenues of research in neurological disorders with a known involvement of BBB disruption. Regeneration of the barrier with existing developmental pathways or the yet unidentified interplay thereof could prove to be an interesting field for future therapeutic strategies. Continued efforts to understand the fundamental mechanisms behind BBB development and maintenance will widen our view on the possibilities to redefine or restore the barrier upon disruption, and is therefore a crucial part of disease-related BBB research.

**Disclosure of Potential Conflicts of Interest**

No potential conflicts of interest were disclosed.

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