3D Culture and Characterization of Blood-Brain Barrier Endothelial Cells in a New Microfluidic Platform

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Research

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Abstract

**Background:** The blood-brain barrier (BBB) protects the central nervous system (CNS) from harmful substances in the peripheral blood stream and ensures brain homeostasis by regulating the transport of nutrients and metabolites. It also prevents many pharmaceutical drugs from reaching their target site and dysfunctions of the BBB are often associated with CNS diseases.

In order to provide a new *in vitro* platform that allows the investigation of barrier characteristics of the BBB and drug transport under physiological and pathological conditions, the TransBBB system was developed. The microfluidic chip consists of ten parallel culture chambers, each allowing the cultivation of endothelial cells on a physiological matrix under perfusion.

**Methods:** Porcine brain capillary endothelial cells (PBCECs) were cultivated on the surface of a hydrogel matrix resulting in a barrier which separates the culture chambers into two compartments. A protocol was established in order to obtain a tight cell layer under perfusion and the effect of different perfusion volumes on BBB formation was investigated. The barrier was then characterized in terms of cell viability, barrier integrity and expression of tight junction proteins.

**Results:** PBCECs formed a tight cell layer on the hydrogel surface preventing FITC dextran from permeating the cellular barrier. The presence of ZO-1 at the cell borders was shown indicating a functional BBB. The perfusion experiments suggested a critical role of medium volume for cell layer integrity under perfusion conditions.

**Conclusions:** The TransBBB chip presented here provides a promising, scalable microfluidic platform for the investigation of cellular barriers which can be integrated seamlessly into existing cell culture and drug testing workflows.

**Background**

The blood-brain barrier (BBB) is a highly selective cellular structure separating the central nervous system (CNS) from the peripheral blood stream. The so-called neurovascular unit (NVU) is composed of endothelial cells attached to a basal lamina, pericytes, parenchymal cells like astrocytes and extracellular matrix components [1, 2]. Endothelial cells are connected by tight junction proteins like claudins or occludin preventing paracellular diffusion and resulting in high transendothelial electrical resistance (TEER) [3]. Furthermore, the endothelium expresses various efflux transporters like P-glycoprotein (P-gp), multidrug-resistance proteins (MRP) or breast-cancer-resistance proteins (BCRP) impeding transcellular permeation [4]. As a result, the blood-brain barrier upholds brain homeostasis and prevents possible harmful substances from reaching the brain, but also poses an obstacle for pharmaceutical agents intended for the treatment of CNS diseases [5]. Moreover, many diseases of the CNS are associated with dysfunctions of the BBB, e.g. Alzheimer's disease, Parkinson's disease etc. [6, 7].
Since many features of the BBB are difficult to assess in vivo, capable in vitro models are essential for the development of new drug candidates as well as the investigation of pathological conditions.

Within the scope of preclinical research, potential drug candidates are investigated in cell culture models and animals prior to their application to humans. Although animal models offer a complex organic environment, therapeutic effects as well as possible toxic side effects cannot be predicted sufficiently by animal experiments due to species related differences [8]. Furthermore, the ethical questions regarding the extensive use of animal trials have gained more and more attention in recent years.

The most common cell culture model for the investigation of physiological barriers is the Transwell system [9]. It is a well plate whose wells are divided into two chambers by a membrane. Endothelial cells are cultivated on the membrane allowing the investigation of transport processes across the cellular barrier. While it is easy to handle, suitable for high throughput and enables the possibility of using human cells, it lacks essential physiological features such as a physiological extracellular matrix or flow related effects like shear stress.

To overcome the shortcomings of traditional cell culture models like the Transwell system, several new microfluidic in vitro models, so-called organ-on-a-chip models, have been developed in recent years [10–13]. The vision is the simulation of the smallest functional units of organs in order to investigate physiological and pathological conditions as well as drug transport mechanisms [14, 15]. An ideal organ-on-a-chip model should address the following requirements:

- standard format for seamless integration into existing workflows
- biocompatible, transparent material
- multiple culture chambers for high throughput assays
- integration of perfusion
- physiological extracellular matrix
- option for co-culture experiments
- real-time read-outs
- simple handling

Although remarkable progress has been made in this field in recent years, there is still a need for a system that meets all requirements previously mentioned. Many of the published microfluidic devices are made of polydimethylsiloxane (PDMS) due to its advantages for microfabrication, however, it is also known for the adsorption of hydrophobic compounds [16–20]. Other systems use artificial membranes for the cultivation of endothelial cells rather than a more physiological extracellular matrix [18–23]. Another disadvantage of some devices is the fact that they are connected to tubes and pumps which is complex to handle and therefore susceptible for disruptions.

In this study, a new microphysiological chip made of cyclic olefin polymer (COP) in microwell format produced by microinjection-molding was used to establish a blood-brain barrier model with brain capillary
endothelial cells. One chip contains ten culture chambers and each culture chamber consists of two
channels separated by an array of micropillars. One of the channels is filled with a biochemically
functionalized hydrogel which forms an interface between two pillars. Endothelial cells are seeded on the
hydrogel surface, thus forming a barrier between the two channels. The channel filled with hydrogel
represents the abluminal and the other channel filled with medium the luminal side. The luminal channel
can be perfused with cell culture medium by gravity driven flow induced by a difference of the medium
amount in in- and outlet reservoir. By using a programmable rocker that tilts the chip at specific intervals
at a certain angle, bidirectional perfusion can be controlled over long periods of time without the need of
a complex periphery involving tubes and pumps.

A more comprehensive description of the technical details of the chip will be published elsewhere.

The aim of this study was to demonstrate the suitability and potential of the TransBBB system as a new
_in vitro_ model of the blood-brain barrier.

**Methods**

**Cell culture**

Primary porcine brain capillary endothelial cells (PBCECs) were isolated as previously described [24] and
stored in liquid nitrogen. Brains were obtained by the slaughterhouse Mannheim free of charge. For every
experiment freshly thawed PBCECs were used.

**Preparation of the Chips**

One channel of each culture chamber of the microfluidic TransBBB chip (microfluidic ChipShop GmbH,
Jena, Germany) was filled with a mixture of the components of 3-D Life PVA-CD Hydrogel SG (Cellendes
GmbH, Reutlingen, Germany) and Matrigel (Corning, New York, USA). Matrigel was added to enhance cell
adhesion. Since Matrigel becomes solid at temperatures above 8 °C, all solutions and materials including
the chip were pre-cooled overnight at -20 °C. The additional movie file shows the gel filling procedure in
detail (see Additional file 1). 30 minutes after gel filling the luminal channel was filled with cell culture
medium (Earl's medium M199 supplemented with L-glutamin, Pen/Strep, Kanamycin and fetal bovine
serum (FBS), Biochrom, Berlin, Germany). The chip was placed in the incubator (37 °C, 5% CO_2_) for 24 h
to equilibrate the hydrogel in order to avoid the formation of air bubbles in the microfluidic channels.
PBCECs in equilibrated medium were seeded at a density of 1,5 × 10^7 cells/mL. After a 5 h attachment
period with the chip positioned on its short edge to allow the cells to sediment on the hydrogel surface,
reservoirs were placed on in- and outlet. 60 µL medium was filled in the inlet reservoir and 20 µL extracted
from the outlet reservoir in order to remove unattached cell debris. After 24 h of incubation, perfusion was
started by placing the chip on a programmable rocker (Mimetas GmbH, Leyden, Netherlands). It slowly
tilted the chip (0,1 cycle/min) with −18 ° to +18 ° tilting angle resulting in gravity driven, bidirectional
perfusion.

**Perfusion volume assay**


Preliminary experiments showed a high sensitivity of PBCECs when perfused with 1 mL of cell culture medium. Hence, the influence of the total volume of medium used for perfusion was investigated. Control channels were connected to 1 mL pipette tips as medium reservoir and filled with 1 mL medium. 10 µL tips were used for low volume channels and filled with 20 µL medium. Perfusion was induced by tilting the chip every 10 min. The development of cell morphology was documented over a period of 30 h. Medium was changed after 24 h.

**Viability**
Calcein-Acetoxymethylester (calcein-AM) is a lipophilic, nonfluorescent compound which can be used to visualize viable cells. It is able to penetrate cell membranes where it is cleaved by unspecific intracellular esterases. The resulting calcein is hydrophilic and accumulates in the cytoplasm. In contrast to calcein-AM, calcein is highly fluorescent and can be visualized with a fluorescence microscope.

Cells were washed by removing medium from the reservoirs, adding 60 µL PBS to the inlet reservoir and waiting for PBS to pass the channel. After removing PBS from the reservoirs, cells were stained by adding Calcein-AM solution in medium (4 µg/mL) to the reservoir. After an incubation period of 30 min at 37 °C cells were washed again and imaged with a Leica TCS SP5 II confocal microscope (CLSM).

**Barrier integrity – Permeability**
The integrity of cell layers can be assessed by perfusing the luminal channel with a fluorescent marker that cannot penetrate a confluent cell barrier and observing its permeation into the abluminal channel over time.

Cell culture medium was removed from the reservoirs and replaced with a solution of FITC dextran (20 kDa, 0,5 mg/mL, Sigma Aldrich, Germany). The permeation was documented by taking one picture per minute over 10 min with Leica CLSM. To quantify the permeation rate of FITC dextran into the hydrogel matrix, two regions of interest (ROI) were selected, one on the luminal side of a cell barrier and the other on the abluminal side. The fluorescence intensity of the luminal channel was related to the fluorescence intensity of the abluminal channel using the analyzing software FiJi. The relative fluorescence intensity was plotted over time using Microsoft Excel.

**Barrier characterization – Immunostaining of ZO-1**
The expression of various tight junction proteins is a pivotal characteristic of blood-brain barrier endothelial cells. Zonula occludens-1 (ZO-1) is an important tight junction protein connecting occludin and claudins to the cytoskeleton [25].

After carefully washing the cells with PBS, the reservoirs were removed. 10% neutral buffered formalin (NBF) was filled into the luminal channel and incubated for 10 min at room temperature to fix the cells. Blocking solution consisting of 1% goat serum and 0,25% TritonX in PBS was added and incubated for 1 h in order to permeabilize cell membranes and block unspecific binding sites. Primary anti-ZO-1 antibody with 0,5% BSA and 0,125% TritonX was incubated for 24 h at 4 °C with the channels covered with parafilm to prevent evaporation of the antibody solution. Secondary antibody solution (goat, anti-
mouse AlexaFluor 488) was incubated for 1 h at room temperature. Cells were washed after each incubation step with PBS and imaged with Leica TCS SP5 II confocal microscope.

**Results**

**Culture of PBCECs on TransBBB chip**

The result of the filling of the chip with hydrogel is shown in Fig. 2a. In order to improve visibility, the hydrogel was dyed with methylene blue. The hydrogel forms a smooth surface between pillars providing a culture area for brain endothelial cells. The seeding density of $1.5 \times 10^7$ cells/mL is the highest possible density before clogging of the channels occurs. It was chosen in order to decrease the chance for uncovered hydrogel areas.

**Perfusion volume assay**

The morphology of cells perfused with 1 mL medium was compared to that of cells perfused with 20 µL. Since the hydrostatic pressure only depends on height (h), fluid density ($\rho$) and gravitational acceleration (g) but not on volume (V), the perfusion rate was the same in control and low volume channels. As shown in Fig. 2c, cells perfused with 1 mL detached and perished shortly after starting perfusion. In contrast, cells perfused with 20 µL remained attached and developed over time.

**Calcein-AM staining**

Calcein-AM was used to visualize viability and cell growth of PBCECs in the microfluidic channels of the TransBBB chip 4 days after seeding. As shown in Fig. 2b, cells grew on hydrogel surface as well as on pillars and channel walls. The side view was generated with CLSM by digitally assembling individual focal planes. It reveals that cells grew confluenty on hydrogel surfaces while growth on pillars was less dense.

**Permeability**

FITC dextran 20 kDa is a commonly used fluorescence marker that shows confluence of cell layers and makes any leaks visible. Figure 3c shows the permeation of FITC dextran into the hydrogel matrix in a control channel without cells, in a channel with a leaky barrier and across a tight barrier after 10 min. Fluorescence intensity in the abluminal channel was considerably lower if a confluent cell layer grows on the hydrogel surface. In order to quantify the permeation of FITC dextran, the ratio of the fluorescence intensity in the abluminal channel and the luminal channel over time is shown in Fig. 3d. Channels with confluent cell layers showed significantly lower values than the cell free control channels indicating a tight barrier. Figure 3d also shows how leaky barriers lead to higher permeation rates.

**Immunostaining of ZO-1**

The expression of tight junction proteins is characteristic for brain capillary endothelial cells. The detection of tight junctions is therefore important for the evaluation of blood-brain barrier *in vitro* models. The result of the immunostaining of ZO-1 is shown in Fig. 3ab. A high fluorescence intensity can be seen on the hydrogel surface as the signals of several adjacent cells overlap. The detailed view of cells
growing on the channel wall reveals the localization of the fluorescence signal. It shows that ZO-1 is located at the cellular borders as well as around the nuclei.

**Discussion**

The great demand for new *in vitro* models of the blood-brain barrier has led to the development of various interesting approaches and a remarkable progress in recent years. Microfluidic systems which are able to realistically simulate the smallest functional units of organs have the potential to revolutionize basic research as well as the development of new pharmaceutical drugs and carrier systems. This study introduces a new microfluidic *in vitro* model of the blood-brain barrier.

The TransBBB chip offers a platform with ten parallel culture chambers in microwell format which allows the seamless integration into existing cell culture workflows. Since it is made of COP, it can be produced by injection molding which makes it easy to manufacture on a large scale [26]. Compared to the frequently used PDMS, it also has the advantage of not exhibiting any adsorbing effects for hydrophobic substances [27]. This is particularly important considering the fact that most CNS drugs are relatively hydrophobic [28]. It does not use an artificial membrane as culture area and instead relies on a hydrogel that mimics the physiological extracellular matrix. 3D Life biomimetic hydrogels from Cellendes® have been shown to successfully mimic extracellular matrices [29–31]. Including Matrigel as one of the gel components promotes cell adhesion and differentiation [32, 33]. When filled into the chips with a small micropipette, it forms smooth surfaces between the pillars, offering physiological culture areas.

Formation of air bubbles poses a significant issue for microfluidic systems as they can impede perfusion and also impair cell growth [34]. It is important to know that the solubility of gases in liquids decreases with increasing temperatures. Therefore, cell culture medium used in this study was always equilibrated for at least 24 h in the incubator at 37 °C.

Perfusion is an important advantage of microfluidic systems compared to static models as it has been shown that the resulting shear stress on endothelial cells is a relevant factor for cell differentiation and the expression of tight junctions and transporters [35–37]. The TransBBB chip can be perfused by gravity driven flow induced by a programmable rocker without the use of complex tubing systems. This also allows perfusion with significantly lower medium volumes. The resulting shear stress on endothelial cells was calculated to be in a range from 0,05–0,1 Pa, which is lower than shear rates found *in vivo* (0,3–2 Pa) [37, 38].

The comparison of cell morphology under perfusion with large and small volumes suggests the crucial role of the ratio of total volume to culture area. The relevance of the amount of cell culture medium used in microfluidic systems in general was already discussed by Walker et al. [34] and Sung et al. [39]. So far there is no conclusive explanation for the finding in our experiment but it will be subject of further investigations. However, the results of this experiment underline that it is an important aspect that should be taken into account when developing new microfluidic *in vitro* models.
Cells with the greatest predictive value for blood-brain barrier models would be primary human brain capillary endothelial cells. Due to the difficulty of obtaining primary human cells, PBCECs were used for this study instead. It has already been shown that these have a greater relevance than immortalized cell lines in terms of barrier tightness and protein expression [40–43]. The field of induced pluripotent stem cells (iPSCs) has made a remarkable progress in recent years. iPSCs have the potential to show similar properties compared to primary cells while being readily available [44]. The use of iPSCs in the presented chip offers a promising option for the future.

PBCECs grew preferentially on the gel surface where they had a flat and stretched morphology leading to a dense layer that separated luminal and abluminal channel. The detection of ZO1 with an immunostaining demonstrates the formation of tight junctions. ZO-1 was also detected around the cell nuclei which could indicate a premature stage of cell development [45].

The permeation assay with the commonly used FITC dextran 20 kDa as fluorescence marker enables the assessment of confluency and barrier integrity. The performed determination of the ratio of abluminal and luminal fluorescence intensity allows the evaluation of permeation rates [46].

Moreover, the TransBBB system offers integrated electrodes for in situ measurements of trans-endothelial electrical resistances (TEER). Up to now, no significant TEER values could be measured with PBCECs cultivated in the chip. It has been reported that even minimal gaps in cell layers could lead to drastically reduced TEER values [47]. Since one culture channel of the TransBBB chip contains up to 21 hydrogel surfaces, the probability is quite high that there could be a small gap in the cellular barrier on one of these surfaces. Therefore, the length of the individual channels and thus the number of cultivation areas per channel will be reduced in further developments. This will also allow even more culture chambers to be incorporated into one chip, which would enable greater throughput.

3-D Life biomimetic hydrogel contains peptide sequences which are cleavable by matrix-metalloproteases allowing cellular migration. The use of other cell types of the NVU like peri- or astrocytes is thus possible in order to further improve the relevance of the model.

**Conclusion**

This study presents a new approach of emulating the blood-brain barrier in vitro. This approach allows the cultivation of endothelial cells on a physiologically relevant matrix and perfusion without the need for a tubing periphery. It is built in microwell format and contains ten culture chambers allowing the integration in existing cell culture workflows and parallelization of experiments. It is planned to arrange the fluid ports in 96 well format which will further enhance the usability. The TransBBB system is suitable for the investigation of transport processes of new pharmaceutical drugs and carrier systems as well as the functionality of the BBB under pathological conditions like hypoxia or hyperglycemia and CNS diseases associated with dysfunctions of the BBB like Alzheimer's or Parkinson's disease. Apart from the blood-brain barrier, other relevant barriers such as gut- or kidney barrier could be emulated.
The observation that PBCECs only survive when perfused with small medium volumes is a very interesting finding which will be further investigated in order to obtain a better understanding of the microenvironment in the microfluidic channels.

The discussed shortcomings will be addressed in future developments. Furthermore, the integration of other sensors for \textit{in situ} monitoring of biorelevant parameters is planned.

All in all, the TransBBB provides a promising platform for the emulation and investigation of physiological barriers and it is a good starting point for further developments.

**List Of Abbreviations**

BBB: blood-brain barrier, CNS: central nervous system, NVU: neurovascular unit, TEER: transendothelial electrical resistance, P-gp: P-glycoprotein, MRP: multidrug resistance protein, BCRP: breast cancer resistance protein, PDMS: polydimethylsiloxane, COP: cyclic olefin polymer, PBCEC: porcine brain capillary endothelial cells, PVA: polyvinyl alcohol, SG: slow gelling, FBS: fetal bovine serum, Calcein-AM: calcein-acetoxymethylester, PBS: phosphate buffered saline, CLSM: confocal laser scanning microscopy, FITC: fluorescein isothiocyanate, ROI: region of interest, ZO-1: zonula occludens-1, NBS: neutral buffered formalin, iPSC: induced pluripotent stem cells

**Declarations**

**Ethics approval and consent to participate**

Not applicable

**Consent for publication**

Not applicable

**Availability of data and materials**

The datasets used and analysed during the current study are available from the corresponding author on reasonable request

**Competing interests**

The authors declare that they have no competing interests

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**Author's contributions**
Authors S.T., P.J., M.S. and G.F. designed the study. Authors S.T., F.S. and L.A. performed experiments and data analysis. Authors A.M., P.J., M.S., and G.F. supervised the research. Authors S.T., A.M., P.J., M.S. and G.F. wrote the paper.

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