African Trypanosome-Induced Blood–Brain Barrier Dysfunction under Shear Stress May Not Require ERK Activation

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Abstract

African trypanosomes are tsetse fly transmitted protozoan parasites responsible for human African trypanosomiasis, a disease characterized by a plethora of neurological symptoms and death. How the parasites under microvascular shear stress (SS) flow conditions in the brain cross the blood–brain barrier (BBB) is not known. In vitro studies using static models comprised of human brain microvascular endothelial cells (BMEC) show that BBB activation and crossing by trypanosomes requires the orchestration of parasite cysteine proteases and host calcium-mediated cell signaling. Here, we examine BMEC barrier function and the activation of extracellular signal-regulated kinase (ERK)1/2 and ERK5, mitogen-activated protein kinase family regulators of microvascular permeability, under static and laminar SS flow and in the context of trypanosome infection. Confluent human BMEC were cultured in electric cell-substrate impedance sensing (ECIS) and parallel-plate glass slide chambers. The human BMEC were exposed to 2 or 14 dyn/cm² SS in the presence or absence of trypanosomes. Real-time changes in transendothelial electrical resistance (TEER) were monitored and phosphorylation of ERK1/2 and ERK5 analyzed by immunoblot assay. After reaching confluence under static conditions human BMEC TEER was found to rapidly increase when exposed to 2 dyn/cm² SS, a condition that mimics SS in brain postcapillary venules. Addition of African trypanosomes caused a rapid drop in human BMEC TEER. Increasing SS to 14 dyn/cm², a condition mimicking SS in brain capillaries, led to a transient increase in TEER in both control and infected human BMEC. However, no differences in ERK1/2 and ERK5 activation were found under any condition tested. African trypanosomiasis alters BBB permeability under low shear conditions through an ERK1/2 and ERK5 independent pathway.

Keywords
► African trypanosomes
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African trypanosomes are tsetse fly transmitted pathogens of humans that play an important, and sometimes devastating, role in the health and welfare of people throughout Sub-Saharan Africa. Two subspecies of Trypanosoma brucei, T. b. rhodesiense and T. b. gambiense, cause human African trypanosomiasis (HAT; aka sleeping sickness). Death is inevitable if a patient is untreated. African trypanosomes replicate at the tsetse fly bite site before disseminating from the skin through the hemolymphatic system to infect various organ systems (stage 1). The parasites that cross the blood–brain barrier (BBB) produce central nervous system (CNS) disease (stage 2). Once CNS disease is established, the parasites are shielded from most trypanocidal drugs, the majority of which do not cross the BBB. Indeed, the brain is probably the source for relapses. Recent intravital and vibratome brain imaging studies in the mouse by us and others have demonstrated that the parasites enter the meninges and the parenchyma of the cerebral cortex and cerebellum soon after infection. In fact, some stage-1 drugs may kill trypanosomes localized in the meninges. Neurological features recently reported in T. b. rhodesiense sleeping sickness patients with stage-1 disease provide further clinical support for the potential for early brain entry.

The BBB, comprised brain microvascular endothelial cells (BMEC), separates the brain interstitial fluid from the blood, maintains the stable environment that allows for effective neuronal function, and protects neurons from circulating neurotransmitters, toxic substances, and pathogens. Shear stress (SS) plays an important, dynamic role in regulating the structure and function of BMEC involved in BBB signaling and ultimately neurological function. When exposed to low SS conditions found in brain postcapillary venules (PCV) and high SS found in brain capillaries, BMEC in vitro, respectively, develop low and high transendothelial electrical resistances (TEERs) that are characteristic of brain PCV and capillaries, respectively. This is important because trypanosomes accumulate and cross the BBB at the level of PCVs but not of capillaries.

Consistent with a possible function in the extracellular milieu, T. brucei C1 (papain) family cysteine proteases cathepsin B-like and cathepsin L-like (TbCatL) enzymes retain activity and stability at physiological pH and are partially resistant to serum-derived inhibitors. Using established static in vitro BBB models based on human BMEC, our studies lead to the hypothesis that mobile trypanosomes and cysteine proteases secreted by the parasites can trigger protease-activated receptors (PARs). Working alone, or in concert with other BMEC receptors/ion channels, PARs induce rises of intracellular Ca\(^{2+}\) ([Ca\(^{2+}\)]\(_i\)) within BMECs. These Ca\(^{2+}\) events permit trypanosomes to cross the BBB endothelium paracellularly and initiate precipitous inflammatory events that are the signature of CNS HAT. In cardiomyocytes TbCatL has been shown to increase the propensity for spontaneous [Ca\(^{2+}\)]\(_i\) release (Ca\(^{2+}\) waves) via ryanodine receptors. Overall, trypanosome disruption of Ca\(^{2+}\) handling by host cells defines a common pathogenic mechanism used by the parasites and might identify potential therapeutic HAT targets.

Previous studies with cultured cells have revealed the involvement of extracellular signal-regulated kinase (ERK) 5 and ERK1/2 in many cellular responses, including cell proliferation. Recent genetic studies have identified that ERK5 is essential for cardiovascular development and neural differentiation, whereas ERK1/2 is important for mesoderm formation. ERK5 is ubiquitously expressed in numerous tissues and is activated by a variety of extracellular stimuli, such as cellular stresses and growth factors, to regulate processes such as cell proliferation and differentiation. The requirement of ERK5 in the maintenance of vascular integrity is highlighted by the fact that induced ablation of ERK5 in adult mice is lethal within 2 to 3 weeks as blood vessels become leaky due to EC apoptosis.

To achieve a proper understanding of how trypanosomes and their proteases trigger BBB and neurological dysfunction, we initiated studies that use BBB models that might mimic the dynamic fluid conditions in the brain during an active trypanosome infection. Using electric cell-substrate impedance sensing (ECIS) and a parallel plate human BMEC-based models under SS conditions found in brain PCVs and capillaries, we hypothesize that the trypanosomes would (1) increase BMEC barrier permeability precursory to paracellular crossing and (2) that barrier permeability will be regulated, in part, via phosphorylation of ERK1/2 and/or ERK5, members of the mitogen-activated protein kinase (MAPK) family containing the Thr-Glu-Try activation motif.

**Methods**

**Human BMEC and Trypanosomes**

Human BMEC (≤ passage 13) were maintained as previously described in medium M199 (Lonza, Walkersville, MD) containing 10% fetal bovine serum (Gemini Bio-Products, West Sacramento, CA). The T. b. rhodesiense used is a human cerebrospinal fluid and bloodstream form isolated from a Kenyan patient with sleeping sickness. This parasite was formerly classified as T. b. gambiense but has been reclassified as T. b. rhodesiense based on the presence of the SRA gene. The trypanosomes were maintained in culture in HMI-9 medium.

**Mechanical Stress Exposure**

**ECIS System**

Human BMEC were seeded on fibronectin (BD Biosciences, Bedford, MA) coated 1F8 × 10E ECIS flow arrays (Applied Biophysics, Troy, NY). Although each array can only be used to monitor a single condition (e.g., with or without parasites; low or high SS), each of eight electrode groups will give an independent TEER measurement. After placing in the ECIS station, human BMEC were exposed to continuous unidirectional laminar flow (LF) by driving the perfusion medium with a computerized peristaltic pump (Cole-Parmer, Vernon Hills, IL) connected to a vented 25 mL reservoir and TEER monitored using an ECIS-Z0 instrument (► Fig. 1A).
Parallel-Plate Flow System

Human BMEC seeded on fibronectin coated glass slides (75 × 38 mm; Fisher Scientific, Pittsburgh, PA) that were placed in a parallel-plate flow chamber system (Cytodyne, Sand Diego, CA) were exposed to LF driven by the hydrostatic pressure created by the vertical distance between the reservoirs and the flow chambers (►Fig. 1B).

For both systems, the cells were kept at 37°C in a humidified 95% air/5% CO2 environment. The magnitude of SS was calculated as previously described.31,32 SSs of 2 or 14 dyn/cm² were applied as flow.

Immunoblotting

After experimentation, human BMEC on the parallel plate flow system were washed in ice-cold PBS twice, scraped in RIPA lysis buffer 1X (Millipore, CA) and placed in an ice bath for 45 minutes. Cell extracts were sonicated and centrifuged at 15,000 g for 20 minutes at 4°C, and the supernatant was collected. Equal amounts of protein (30 μg per lane, BioRad protein assay system; BioRad Laboratories Inc, Hercules, CA) were separated by 10% sodium dodecylsulfate-polyacrylamide gel electrophoresis and transferred to a polyvinylidene fluoride membrane (BioRad Laboratories Inc). After blocking for 1 hour with tris-buffered saline containing 0.1% Tween 20 and 5% nonfat dry milk, the membrane was probed with primary antibody, either phospho-anti-ERK5, total anti-ERK5, phosphor anti-ERK1/2, or total anti-ERK1/2 antibody (Cell Signaling, Beverly, MA), and horseradish peroxidase-conjugated antirabbit polyclonal secondary antibody (Cell Signaling), before detection of immunoreactivity by enhanced chemiluminescence (Amersham). All blots were quantified with densitometry (BioImage, Ann Arbor, MI).

Real-Time TEER Measurements by ECIS

After human BMEC confluence on the ECIS flow array was reached under static conditions, 2 dyn/cm² LF was applied to the human BMEC until stable TEERs under SS were again reached. A small volume of concentrated trypanosomes in medium was then added to one chamber so that a final parasite concentration of 5 × 10⁶ parasites/mL was attained. Medium without parasites was added to the other array and used as the sham control and TEER measurements taken every 30 seconds. At a later time point, LF was increased to 14 dyn/cm² and TEERs recorded.

Results

We have demonstrated that different types of SS regimens have different effects on ECs and may account for the variable response of ECs to hemodynamics in the circulation.30 ►Fig. 2 shows that the initial TEER values for the static confluent human BMEC monolayers were approximately 1,000 Ω. Upon application of 2 dyn/cm² of LF to the ECIS flow arrays, TEER values increased approximately 60% and plateaued in about an hour. Addition of the trypanosomes under these SS conditions lead to a rapid 15% drop in BMEC TEER. Furthermore, when SS was increased to 14 dyn/cm², transient increases in BMEC TEER were observed: the control BMEC responded by initially getting tighter, followed by a drop in TEER going back to the initial TEER values under the
lower SS (2 dyn/cm²) condition. Remarkably, TEER values for the BMEC infected with trypanosomes also rapidly increased during this same time period and eventually reached control TEER values.

Phosphorylation of ERK1/2 and ERK5 were used as a measure of their activation. – Fig. 3 shows that when human BMEC were exposed to LF at 2 and 14 dyn/cm² for 4 hours, P-ERK5 did not significantly increase under any of the flow conditions in the presence or absence of trypanosomes. ERK1/2 levels also did not show they were affected by the parasites but may have been affected by different types of flow and SS. These findings support a previous study showing trypanosome crossing of human BMEC grown on Transwell (Corning, Tewksbury, MA) inserts is impaired by inhibitors of PI3K (phosphatidylinositol 3-kinase) and p38 MAPK, but not ERK1/2.34

Discussion

Our results demonstrate that African trypanosomes are able to induce BBB opening precursory to trypanosome transmigration into the brain under SS conditions found at PCVs, a known BBB entry site for the parasites. Our study also lend support to recent in vivo intravital and vibratome brain imaging studies in mice by us and others that show that African trypanosomes can enter the brain soon after entering the systemic circulation.

The results from the Western blot analysis reveal an increase in ERK1/2 and ERK5 activation with the addition of flow. Under static conditions, the addition of trypanosomes activated both ERK1/2 and ERK5. Similarly, exposure of BMECs to 2 and 14 dyn/cm² of LF also showed elevated levels of phosphorylated ERKs. Under LF conditions, there appears to be an increase in expression levels in BMEC as SS under low PCV conditions (a putatively permissive BBB crossing site) is increased to the higher high SS conditions found in brain capillaries, a less permissive crossing site. Addition of trypanosomes did not change this profile.
While ERK5 was activated, our results indicate that activation by flow and trypanosomes are not additive processes. Confluent BMECs exposed to trypanosomes had similar levels of activation as those exposed to high and low SS. However, both the presence of flow and trypanosomes did not further elevate levels of P-ERK1/2 and P-ERK5. These data are in concurrence with our previous data using ERK5 silencing in BMECs. Addition of these knock out genes did not prevent a concurrence with our previous data using ERK5 silencing in elevate levels of P-ERK1/2 and P-ERK5. These data are in both the presence of activation as those exposed to high and low SS. However, changes in human BMEC permeability. Our findings demonstrate that African trypanosomes cause changes in BBB permeability through an ERK1/2 and ERK5 independent pathway. Future studies to assess the role of other putative signal molecules that regulate permeability are warranted and will be facilitated using this in vitro model.

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Conflict of Interest
The authors have declared that no conflict of interest exists.

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