Equid Herpesvirus-1 Exploits the Extracellular Matrix of Mononuclear Cells to Ensure Transport to Target Cells

HIGHLIGHTS

Several viral proteins facilitate rapid and efficient virus cell-to-cell spread

Viruses can evade face-off engagement with host defenses via cell-to-cell spread

The ECM of PBMC ensures safe transmission of viruses to target cells

Viruses can be transmitted during a transendothelial migration process of PBMC
Equid Herpesvirus-1 Exploits the Extracellular Matrix of Mononuclear Cells to Ensure Transport to Target Cells

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SUMMARY

Mononuclear cells are the first line of defense against microbial infection. Yet, several viruses have evolved different mechanisms to overcome host defenses to ensure their spread. Here, we show unique mechanisms of how equid herpesvirus-1 manipulates peripheral blood mononuclear cells (PBMC) to travel further in the body. (1) “PBMC-hitching”: at the initial contact, herpesviruses lurk in the extracellular matrix (ECM) of PBMC without entering the cells. The virus exploits the components of the ECM to bind, transport, and then egress to infect other cells. (2) “Intracellular delivery”: transendothelial migration is a physiological mechanism where mononuclear cells can transmigrate through the endothelial cells. The virus was intangible and probably did not interfere with such a mechanism where the infected PBMC can probably deliver the virus inside the endothelium. (3) “Classical-fusion”: this process is well mastered by herpesviruses due to a set of envelope glycoproteins that facilitate cell-cell fusion and virus spread.

INTRODUCTION

Cell-to-cell transmission is one of the strategies adopted by several viruses to overcome host barriers and to establish successful infection (Zhong et al., 2013). Several advantages have been documented for this mode of transmission that center mainly on avoiding face-off engagement with host defenses (Gupta et al., 1989; Mothes et al., 2010; Sattentau, 2008). Viruses have evolved several mechanisms, including cell-cell fusion, synapse formation, and induction of actin tails, filopodial bridges, or nanotubes, and are equipped with the necessary information to facilitate a secure and rapid cell-to-cell dissemination (Aubert et al., 2009; Bennett et al., 2009; Sherer et al., 2007; Sowinski et al., 2008). Other modes of cell-to-cell transmission involve the utilization of the carbohydrate-rich extracellular matrix (ECM) (Mothes et al., 2010). The ability of viruses to utilize and hijack cell-cell contact machineries contributes to the success of viral infection and spreading.

The ECM is a complex three-dimensional structure present in all tissues that is generally composed of well-organized networks of macromolecules (including polysaccharides and proteins) with distinct biochemical and biomechanical properties (Stavolone and Lionetti, 2017). Moreover, besides providing a physical support for tissue integrity and elasticity, the ECM contributes functionally to the regulation of cell growth, survival, motility, intercellular adhesion and communication, as well as differentiation (Bonnans et al., 2014; Stavolone and Lionetti, 2017). The ECM is a dynamic structure that is constantly remodeled and specifically tuned in response to different physiological cell conditions as well as in response to diseases and infections (Bonnans et al., 2014; Humphrey et al., 2014). Although ECM represents a formidable barrier against pathogen invasion, many viruses have evolved diverse mechanisms to overcome and even exploit the ECM for entry and spread (Akhtar and Shukla, 2009; DiGiuseppe et al., 2017; Dimitrov, 2004). Human T cell leukemia virus type 1 (HTLV-1) was shown to infect T lymphocytes and then bud into the ECM, which carries the virus through virus-induced ECM components allowing virus spread and infection of other T lymphocytes (Pais-Correia et al., 2010).

Infection of peripheral blood mononuclear cells (PBMC) plays a key role in disease outcomes because PBMC have access to all body tissues. Although non-adherent, PBMC can adhere to endothelial cells (EC) after stimulation, as is presumably the case after certain viral infections, where viruses can attach to,
enter, and replicate in EC (Keller et al., 2003; Valbuena and Walker, 2006). The PBMC-EC interface thus represents a crucial spot that can influence virus pathogenesis and disease outcomes. Infection of EC through infected PBMC has been documented for members of several virus families, including bunya, herpes, flav, toga, and arena viruses (Charrel and de Lamballerie, 2010; Chu et al., 2016; Gowen and Hickerson, 2017; Kunz and de la Torre, 2017; Ma et al., 2013; Mackow and Gavrilovskaya, 2009). However, the mechanism of how these viruses spread from infected PBMC to EC is only poorly understood. Disease outcomes of these viral infections are mostly fatal (to humans and animals) due to hemorrhagic fevers, severe respiratory diseases, as well as neurological and/or congenital disorders.

Herpesviruses represent a large group of DNA viruses that infect a wide variety of hosts. However, the mechanism(s) used by herpesviruses to spread from cell to cell differs greatly depending on virus species and their dedicated target cells or tissues. Cell-cell fusion, synapse formation, filopodial bridges, and nanotubes were documented for cytomegaloviruses (CMV), varicella zoster virus (VZV), pseudorabies virus (PRV), and herpes simplex virus (HSV-1) (Cole and Grose, 2003; Dingwell et al., 1994; Favoreel et al., 2005; Gerna et al., 2000; Sweet, 1999; van Leeuwen et al., 2002). Infection of PBMC with PRV, CMV, and VZV and subsequently their transfer to EC represents a critical step for viral pathogenesis. CMV and PRV were documented to spread between PBMC and EC via cell-cell (micro)fusion (Digel et al., 2006; Gerna et al., 2000; Van de Walle et al., 2003). Equine herpesvirus type 1 (EHV-1) is another example of herpesviruses infecting PBMC and then being transferred to EC via cell-associated viremia causing severe outcomes including abortion, perinatal mortality, neurological disorders, and death (Chowdhury et al., 1986; Van Maanen, 2002). Primary virus replication occurs in the respiratory epithelia of infected horses. The virus rapidly reaches the lymphoid tissue associated with the upper respiratory tract, infects mononuclear cells that enter bloodstream, and carry the virus within a short time frame to the vasculature of tissues, such as the pregnant uterus or the central nervous system. The virus can spread and replicate in EC causing different pathological injuries such as vasculitis, thrombosis, and ischemia (Smith et al., 1996; Wilson, 1997). However, the exact mechanism of virus spread is still unknown. EHV-4 infection, on the other hand, is mainly restricted to the upper respiratory tract with very rare events of leukocyte-associated viremia (Osternieder and Van de Walle, 2010; Vandekerckhove et al., 2011). It is likely that this clear difference in EHV-1 and EHV-4 pathogenesis and disease outcomes is associated with differences in gene products, despite the significant genetic and antigenic similarity, that facilitate virus spread from cell to cell (Sattentau, 2008). Using dynamic in vitro models, we recently showed that EHV-1 was able to maintain tethering and rolling of infected PBMC on EC, which resulted in virus transfer from PBMC to EC (Spiesschaert et al., 2015a). Most remarkably, no EHV-1-productive infection in PBMC was observed, which, however, does not exclude unambiguously restricted productive virus replication albeit at low levels (Drebert et al., 2015; Laval et al., 2015; Spiesschaert et al., 2015a).

Here, we combine confocal imaging, live-cell imaging, and electron microscopy analyses together with functional assays to study virus cell-to-cell spread between PBMC and EC. Our data unravels unique mechanisms of cell-to-cell transmission exploited by herpesviruses, in which the virus is embedded in the ECM of PBMC without entering or infecting the cells. The embedded viruses were protected against circulating neutralizing antibodies until the PBMC reached the EC, where the virus was released to infect the endothelium. We were also able to document several transendothelial migration events of mononuclear cells through EC, where infected PBMC might be able to deliver the virus directly inside the EC.

RESULTS

Virus Embedding in the Carbohydrate-Rich Extracellular Matrix Structures

Confocal microscopy was performed to localize virus particles with respect to the plasma membrane and the ECM of PBMC. We used an EHV-1 strain with a red fluorescent (mRFP) protein fused to the small capsid protein VP26 (EHV-1RFP, to facilitate virus particle tracking) and the fluorescein isothiocyanate (FITC)-labeled plant lectins (ConA and WGA) to stain glycan-rich carbohydrate components of the ECM. EHV-1RFP (multiplicity of infection [MOI] = 0.5) was added to PBMC for different time periods (5 min, 1 h, 24 h, 2 days, 3 days, 5 days, and 7 days) at 37°C, treated with ice-cold citrate buffer (pH 3) for 1.5 min to get rid of ECM-unbound viruses, and then fixed with paraformaldehyde 4%.

Interestingly, we found that virus signals (either single viruses or clusters) were colocalizing with the ECM at all time points, even after 7 days (5 min: Figures 1A and 1B; 1 and 24 h: Figures S1A and S1B; 1–7 days: Figure S2). The 3D image with virus particles colocalizing with the ECM after 5 min (Figure S1D) showed...
embedding of EHV-1 viral particles in these structures. We only detected virus particles inside the infected cells after 24 h of infection and up to 7 days (Figures S1C and S2).

To further confirm that virus particles were embedded in the ECM and not just bound to cell plasma membrane, EHV-1 RFP (MOI = 0.5) was added to PBMC for 5 min at 37°C. The cells were stained with CellVue dye to stain plasma membrane and FITC-labeled ConA to stain ECM. It was clear that virus particles were co-localizing only with the ECM (Figure S3).
Figure 2. Virus Transmission from Infected PBMC to EC under “Static” and “Dynamic” States

(A–F) Schematic depiction of contact (A), transwell (B), and flow chamber assays (C) is shown. PBMC were infected with EHV-1 GFP for different time periods (5 min, 1 h, and 6 h). Infected PBMC were overlaid on EC under “static” conditions (D) or allowed to flow over EC “dynamic” (E and F) in the presence of neutralizing antibodies. Under dynamic conditions, whole PBMC population (E) and/or each PBMC subpopulation (T-lymphocyte, B-lymphocyte, monocyte); (F) were infected for 5 min either at 37°C (E and F) or at 4°C (E). After 24 h, virus spread was assessed by counting the plaques on EC excluding the inlet and outlet of the slide. As a control, infected PBMC or PBMC subpopulation were placed into a transwell insert without direct contact between EC and PBMC “no contact.” The data represent the mean ± standard deviation of three independent and blinded experiments. Significant differences in plaque numbers were seen between different infection points under “static” conditions (*) (n = 3; one-way ANOVA test followed by multiple comparisons tests; *p < 0.05; **p < 0.01). PBMC in (F) refers to whole PBMC population.

(G) Confocal microscopy of overlaid infected PBMC on endothelial cells after 2 h. ECM was stained green with FITC-labeled ConA (green), EHV-1 RFP viral particles are red (arrowhead), and nucleus was stained with DAPI (blue). Data are representative of three independent experiments. Scale bar, 10 μm, and scale bar of magnification, 7 μm. Image stacks (number of stacks = 17 with 0.75 μm z stack step size) were photographed using VisiScope Confocal FRAP microscope. Presented here is a single optical section of the stacks.
Comparison between infected (Figures 1A and 1B) and non-infected cells (Figures S4A and S4B) showed no significant differences with respect to organization of the ECM glycans in infected and non-infected cells (Figure S5).

We next looked at the components of the ECM and their colocalization with EHV-1RFP. Confocal microscopy analyses were carried out after 5 min infection of PBMC with EHV-1RFP (MOI = 0.5). The cells were stained for several surface proteins including collagen, agrin, and ezrin (Pais-Correia et al., 2010). We found that virus particles were colocalized and clustered with collagen and agrin; however, there was no significant reorganization of these proteins among infected and non-infected cells (Figures 1C, 1D, S4C, and S4D). There was almost no colocalization of the virus particles with ezrin with no reorganization (Figures 1E and S4E). Heparinase enzyme treatment that hydrolyzes heparan sulfate (HS) proteoglycans resulted in destruction of the ECM-bearing viral particles (Figure 4C) indicating their involvement as traps and/or carriers of viruses.

Spread of ECM-Associated Viruses from Infected PBMC to Endothelial Cells

To test virus transmission from infected PBMC to EC and the role of extracellular carbohydrate-rich matrix structures, overlay assays (under either static or dynamic conditions) were performed. EHV-1GFP virus (EHV-1 expresses the enhanced green fluorescent protein [EGFP] for rapid identification of infected cells) was added to PBMC for different time periods (5 min, 1 h, or 6 h) followed by citrate treatment as mentioned above. Infected PBMC were then overlaid on EC in “contact” model (Figure 2A) for 2 h in the presence of virus-neutralizing antibodies to prevent the infection of free virus particles, followed by extensive washing several times to get rid of unbound PBMC. The “no-contact” model was used as a control to exclude cell-free virus transmission (Figure 2B). Our data showed EHV-1 transmission from infected PBMC to EC at all time points in the presence of neutralizing antibodies, including the early time point (5 min; Figure 2D). This indicates that ECM-embedded viruses on the surface of PBMC were able to spread and replicate in the target EC without previous entry into PBMC. In all cases, no virus transfer from infected PBMC to EC was observed under “no-contact” conditions at all time points (Figure 2D).

As was shown in the stationary setup, we further expanded our investigation to test EHV-1 transfer under dynamic conditions using a fluidic setup (flow chamber assay; Figure 2C). In this assay, EHV-1GFP was added to PBMC for 5 min (at 37°C or at 4°C). At 4°C, viruses are attached to the cells without entry or penetration. Cells were allowed to flow over the confluent EC monolayer in the presence of virus-neutralizing antibodies to neutralize free virus particles. Our data showed that the 5-min-infected PBMC were able to transfer the virus to EC under flow condition (Figure 2E). To determine which PBMC subpopulation might be responsible for virus spread, PBMC were sorted and different subpopulations (T lymphocytes, B lymphocytes, or monocytes) were infected with EHV-1 for 5 min and the cells were allowed to flow over EC in the presence of virus-neutralizing antibodies. All PBMC subpopulations were able to transfer the virus to EC (Figure 2F). In all cases, no virus transfer from infected PBMC or the subpopulations to EC was observed under “no-contact” conditions (Figures 2E and 2F).

To further visualize virus transfer from PBMC to EC, EHV-1RFP was added to PBMC for 5 min, overlaid on EC for 2 h, and analyzed using confocal microscopy. The z stack immunofluorescence confocal images came in line with the other infection assays showing PBMC in close proximity to EC with viral particles on the surface of infected PBMC and the underneath EC (Figures 2G and S6).

To track EHV-1 transmission from the surface of infected PBMC to EC, time-lapse live cell imaging was performed. After 5 min infection of PBMC with EHV-1RFP+GFP, the cells were added to EC and visualized by time-lapse live cell imaging confocal microscopy. Interestingly, virus movement (represented by red signals) from infected PBMC to EC was tracked at the early time points (within the first hour after adding infected PBMC) and virus replication in EC (represented by GFP expression and production of progeny...
RFP-labeled viruses) was visualized 2–3 h post overlay (Figures 2H and S7 and Video S1). No virus replication (no GFP expression) was detected in PBMC carrying the virus. The free red signals found in the field might represent free virus particles or cell debris. It was clear that free virus particles, if present, were not able to infect EC due to the presence of neutralizing antibodies in the medium. This conclusion was confirmed by the absence of GFP-expressing EC (except the one with PBMC on top) and the transwell control assay (Figures 2D and 2E).

Taking all previous results together, the data display the role of extracellular carbohydrate-rich matrix in virus PBMC-EC transmission that physically protects the virus against virus neutralizing antibodies and possibly other immune reaction. This carbohydrate-rich ECM not only has a role in virus transmission between PBMC and EC but also probably has a role in viral transfer between PBMC themselves as the PBMC tend to aggregate together in the infection than the non-infection state (Spiesschaert et al., 2015a; van der Meulen et al., 2001) (Figure S8).

**Correlative Fluorescent and Electron Microscopy Identifies Virus Particles in the Interface between PBMC and EC**

Compared with fluorescence microscopy, transmission electron microscopy (TEM) provides a significantly higher resolution, however, at the price of a much smaller field of view and elaborate sample preparation. To further elucidate the mechanism of virus transfer between PBMC and EC at ultrastructural level, we applied a correlative workflow to identify and target individual virus-loaded PBMC in close proximity to EC. EC were grown on gridded coverslips, incubated with EHV-1 RFP (MOI = 5) for 5 min and overlaid on EC. Red: mRFP-labeled EHV-1, blue: Hoechst-stained nuclei. The insert shows one PBMC (red arrow) carrying a virus load and attached to one endothelial cell. The dashed line indicates the direction of the targeted plane for ultrathin sectioning. TEM images of targeted PBMC show virus particles (white arrows) on the surface of infected PBMC (B and C) as well as one virus particle (white arrow) in the contact zone between infected PBMC and EC (D–F). Marked parts with rectangles in (B and E) are magnified in (C and F), respectively. Scale bar, 1 μm. See also Figure S9.
based on their position relative to the coverslip grid (Figure 3A). We observed the presence of enveloped EHV-1 particles on the surface of infected PBMC that are approaching the resting EC (Figures 3B, 3C, and S9). Interestingly, virus particles were found in the intercellular space between tightly interacting infected PBMC and EC (Figures 3D–3F). We did not find evidence of virus budding from the PBMC or fusion events between infected PBMC and EC plasma membranes at this time point (5 min infection of PBMC and 1 h overlaying incubation period, see Transparent Methods). This further indicates that virus particles might be transported embedded on the surface of PBMC and do not necessarily need to enter them.

**Destruction of ECM-Associated Virus Clusters Reduces Virus Spread**

To unravel the prominence of the extracellular carbohydrate-rich matrix in virus transmission from PBMC to EC, extracellular viral clusters were disrupted mechanically by vigorous pipetting, chemically (heparin treatment; EHV-1_5M_Cell_hep), or both (EHV-1_5M_Cell_pip_hep). The disrupted cells (A) or the supernatant (EHV-1_5M_SUP_pip, EHV-1_5M_SUP_hep, EHV-1_5M_SUP_pip_hep) (B) were added to EC. As a control, the ECM was left undisturbed (EHV-1_5M_Cell or EHV-1_5M_SUP). The data represent the mean ± standard deviation of three independent and blinded experiments. Significant differences in plaque numbers on EC were seen between the different treatment procedures. n = 3; one-way ANOVA test and Dunnett’s multiple comparisons test; **p < 0.01, ***p < 0.001.

(C and D) Treatment with heparinase III or heparin fractionates and elution of ECM-viral assemblies. Confocal microscopy showing infected PBMC with EHV-1 RFP + GFP (red) treated with heparinase III (C) or heparin (D) and stained for cell surface carbohydrate-rich matrix with ConA (green). PBMC nucleus was stained with DAPI (blue). Data are representative of three independent experiments. Scale bar, 10 μm, and scale bar of magnification, 7 μm. Image stacks (number of stacks = 17 with 0.75 μm z stack step size) were photographed using VisiScope Confocal FRAP microscope. Presented here is a single optical section of the stacks.
and added to EC monolayers. Treatment with heparinase III or heparin leads to the detachment of ECM-viral assemblies (Figures 4C and 4D). Partial destruction of the ECM clearly leads to a significant reduction in virus transfer between PBMC and EC accompanied by reduction of EC infection rates (Figure 4A). On the other hand, supernatants from treated PBMC contained more ECM-viral assemblies and gained a significant increase in infectivity after adding to EC, even in the presence of virus-neutralizing antibodies (Figure 4B). Although PBMC treatment significantly reduced virus spread, it did not completely abolish virus transmission due to the remaining ECM-virus clusters on the surface of PBMC that can still transfer the virus (Figures 4C and 4D). These results indicate that integrity of ECM is essential in virus transmission from PBMC to EC.

Transendothelial Migration of PBMC through the Endothelial Cell
Mononuclear cell migration across the endothelial barrier is a distinct process that can be through either the endothelial junctions (paracellular) or the EC body (transcellular) (Muller, 2013; Schimmel et al., 2017). The process of transendothelial migration is usually preceded by orchestrated steps starting with rolling, firm adhesion, and microvillar projections invaginating the endothelium that might trigger transmigration (Schmidt et al., 2011). While running TEM analysis, we noticed several PBMC behaving as they are preparing for transmigration; they firmly attached to EC and having projections into the EC (Figures 5A and 5B). We further detected intact PBMC transmigrating through the EC (Figures 5C–5F). Interestingly, we could identify in several occasions incorporated infected PBMC (with virus particles) inside EC; however, they seem to be partially digested (Figures 5G–5L and S10). We further detected virus particle on the surface of a digested PBMC within EC (Figures 5G–5I). This pathway might represent a novel strategy and a mechanism of direct intracellular cell-to-cell spread of viruses.

Cell-to-Cell Fusion Events between Infected PBMC and EC
To test the possible incidence of classical fusion events between infected PBMC and EC, PBMC were transfected with mRFP-expressing plasmid and infected with EHV-1\textsuperscript{GFP} for 24 h to allow virus replication and expression of virus fusion machinery (namely, gD, gB, gH-gL) on the surface of infected PBMC (Figure S11). PBMC were then co-cultured with the target EC for 3 h. Upon adhesion, infected PBMC fused with the underlying uninfected EC, which leads to the transfer of their red cytoplasm as well as intracellular virions between the two cells. EC expressed GFP due to virus replication after transfer through the fusion canal (Figure 6A).

In another assay, PBMC were transfected with T7 RNA polymerase-encoding plasmid and then infected with EHV-1\textsuperscript{GFP} (Trans_inf_PBMC). PBMC were overlaid on EC transfected with a plasmid encoding luciferase gene under a T7 promoter (Trans_EC). As negative controls, PBMC were either transfected with the T7 RNA polymerase-encoding plasmid without infection or infected with EHV-1\textsuperscript{GFP} without transfection. In all cases, the EC were transfected with a plasmid encoding luciferase under a T7 promoter. As a positive control, PBMC were transfected with the two plasmids encoding T7 RNA polymerase and luciferase under a T7 promoter. Luciferase activity was recorded as a measure of cell-cell fusion. PBMC-EC fusion events were only detected in the setup where PBMC were first transfected and infected and then overlaid on transfected EC. Transfection of PBMC with both plasmids gave the same luciferase activity (Figure 6B).

No Transmission Events for ECM-Clustered EHV-4
We have shown before that EHV-4 was unable to spread (under contact or dynamic conditions) from infected PBMC to EC (Spiesschaert et al., 2015a). Here, we show that EHV-4, like EHV-1, is clustered with the extracellular carbohydrate-rich matrix of infected PBMC (Figure S12A). However, infected PBMC were not able to transfer EHV-4\textsuperscript{GFP} to EC as shown in the contact assay (Figure S12B) and the flow chamber assay (Figure S12C). Destruction of the extracellular carbohydrate-rich structures from infected PBMC by extensive pipetting and heparin treatment released ECM-virus assemblies into the medium; however, these assemblies were not infectious to EC as shown for EHV-1 (Figure S12D). It is worth mentioning that EC preparations are permissive for EHV-4 infection using free infectious virus produced by equine dermal (ED) cells (Figure S12E).

Viral gB and gD Influence ECM-Associated Virus Cell-to-Cell Transmission
From our results, it was clear that both EHV-1 and EHV-4 are embedded in the ECM of infected PBMC before entering the cell itself. However, only EHV-1 was able to step forward and spread to infect EC at
Figure 5. Transcellular Migration of Infected-PBMC through Endothelial Cells

(A–L) Ultrathin cross sections of PBMC adherent to and crossing endothelial cells. PBMC were infected for 5 min with EHV-1 RFP and incubated over adherent EC for 3 h. Multiple intracellular protrusions invaginating the apical endothelial membrane (A and B; black arrows) and PBMC migrating through the endothelial cells (C) are shown. Intact PBMC is shown within the cytoplasm of EC (D–F; red arrow). Migrating PBMC (E) is marked with a rectangle and further magnified (F, red arrow). Partially digested infected PBMC is shown (G–L) and marked with a rectangle, which is magnified (H–I and K–L). Virus particles attached to migrating PBMC (either outside, H and I, or inside, K and L) are marked with white arrows. Scale bar, 1 μm. See also Figures S10 and S13.
early time point. Several factors can be involved in this privilege of EHV-1; of interest are the envelope glycoproteins involved in virus entry (Azab and Osterrieder, 2017b). The role of two envelope proteins gB and gD in virus transmission was tested. The two genes were swapped between EHV-1 and EHV-4 (Azab and Osterrieder, 2012; Spiesschaert et al., 2015b), PBMC were infected for 5 min with either EHV-1_gB4 and EHV-1_gD4 or EHV-4_gB1 and EHV-4_gD1, and infected PBMC were added to EC in "contact" and "flow chamber" models. Interestingly, swapping the two genes significantly affected ECM-associated virus spread from infected PBMC to EC. The recombinant viruses (EHV-4_gB1 and gD1) were able to spread and infect EC under "contact" condition only (Figures 7B and 7D). On the other hand, EHV-1_gD4 and EHV-1_gB4 showed a significant reduction in virus transfer when compared with parental EHV-1, under both static and dynamic conditions (Figures 7A and 7C).

DISCUSSION
As successful parasites, viruses have evolved several and diverse mechanisms to facilitate rapid cell-to-cell dissemination and to promote immune evasion (Mothes et al., 2010; Sattentau, 2008). Different pathways of cell-to-cell spread have been identified that fall mainly into two main categories; (1) cell-free viruses: where intracellular viruses are released into the extracellular space and infected new cells; (2) direct cell-to-cell contact: where intracellular viruses spread from one cell to another without being extracellularly egressed (Marsh and Helenius, 2006; Mothes et al., 2010; Sattentau, 2011). Here, we document two unique pathways of virus transmission that include (1) the embedding of virus particles in the ECM without entering the cells and (2) a possible intracellular delivery during transendothelial migration of infected PBMC into EC. The ECM-associated viruses are protected from the immune attack and efficiently spread from infected...
PBMC to other immune cells or to EC. Furthermore, infected PBMC can transmigrate through the EC where ECM-associated viruses can be delivered directly inside EC.

The portal of entry is not the ultimate goal of herpesviruses, and they need to travel (through either blood or nerve) to their preferred tissues where they can establish latency or induce disease (Adler et al., 2017). To achieve this goal, different mechanisms of direct cell-to-cell spread have been documented including cell-cell fusion, synapse formation, or tunneling nanotubes (Gerna et al., 2000; Jansens et al., 2017; Saksena et al., 2006). The pathogenesis of EHV-1 is distinct from other members of the Herpesviridae family. Infection starts first at the respiratory epithelium where PBMC contracts the virus and delivers it to the endothelium (Ma et al., 2013; Smith et al., 2001). Infection of PBMC and subsequent virus spread to the endothelium are still to be understood. We and others have shown that virus replication in PBMC is “restricted” without obvious viral protein production or expression on the cell surface. However, when infected PBMC reach the endothelium, the virus finds its way outside the PBMC to infect new cells, probably through cell-cell fusion (Laval et al., 2015; Spiesschaert et al., 2015a). This pathway provides a secure trip for the virus without being attacked by the immune system. In the mechanism that we provide here for the first time, PBMC (during the first hour of infection) carry the virus on its surface associated with the ECM; the virus does not enter the cells. Although appeared to be exposed to the surrounding environment, viruses are fully protected against neutralizing antibodies that prevented free virus spread. Upon contact with EC either under “static” or “dynamic” conditions, the virus can spread and infect the endothelium. We further showed that all three

![Figure 7. Role of Viral gB and gD in VirusTransmission](image)

(A–D) PBMC were infected with the indicated viruses (MOI = 0.1) for 5 min. Infected PBMC were overlaid on EC under “static” conditions (A and B) or allowed to flow over EC “dynamic” (C and D) in the presence of neutralizing antibodies. After 24 h, virus spread was assessed by counting the plaques on EC. As a control, infected PBMC were placed into a transwell insert without direct contact between EC and PBMC “no contact.” The data represent the mean ± standard deviation of three independent and blinded experiments. Significant differences in plaque numbers were seen between the different viruses as indicated. One-way ANOVA test followed by Dunnett’s multiple comparisons test (B) or Kruskal-Wallis test with Dunns multiple comparisons test (A and C); *p < 0.05. See also Figure S12.
subpopulations of PBMC (T lymphocytes, B lymphocytes, and monocytes) could transfer the virus to EC early after infection. It would be interesting, for future studies, to stain ECM on the surface of each subpopulation to determine virus localization on the surface of these cells.

We next asked the question whether this mechanism is spontaneous and any virus can be stuck to the surface of PBMC and then spread. EHV-4, a close relative to EHV-1 and having a restricted pathogenesis to the upper respiratory tract, was not able to spread to the endothelium, although it was also stuck to the ECM. This indicates that EHV-1 exploits the existing natural ECM and hides in this complex network until it reaches the endothelium where it can exit the surrounding matrix and infect EC. ECM-associated viruses can also be spread to other non-infected PBMC in the circulation or in secondary lymphoid organs. It was shown that dendritic cells contract EHV-1 infection from the respiratory epithelium (Vandekerckhove et al., 2011); in turn, other cells may get infected after contacting EHV-1 clusters on the surface of dendritic cells. A similar but distinct mechanism was described before for HTLV-1 virus (Pais-Correia et al., 2010). It has been shown that HTLV-1-infected lymphocytes produce and store viral particles as extracellular assemblies attached to cell surface extracellular components through a virus-induced mechanism. HTLV-1 seems to hijack certain host cell proteins and enhance their expression to build the extracellular assemblies. In that case, virus infection takes place first within the cells where the virus can modify the carbohydrate compositions and/or their spatial organization. The virus then buds into and accumulates in the extracellular structures that may locally increase the infectious titer and facilitate HTLV-1 cell-to-cell spread (Pais-Correia et al., 2010).

In the natural situation where horses are infected with EHV-1, PBMC transfer the virus from the site of primary infection in respiratory epithelia to the endothelium. In this pathway, PBMC might take hours to reach lymph nodes and days until viremia develops. We show here that PBMC can keep the virus on its surface (associated with the ECM) for up to 7 days, which may point to their ability to transfer the extracellular viruses to EC. However, transfer of intracellular virus particles through cell-cell fusion or any other mechanism cannot be excluded.

Interestingly, we detected enveloped EHV-1 in the interface between tightly interacting infected PBMC and EC. Our correlative approach enables us to roughly target infected PBMC in close proximity to EC during TEM sample preparation, whereas visualizing virus particles with ~180 nm diameter within 50- to 70-nm ultrathin section through an entire cell is still a rare event and thus remains challenging. Fluorescence signals from single labeled virus particles are weak, and most fluorophores are not compatible with resin-embedding procedures. Using fluorescence imaging is thus difficult to target rare events on ultrathin sections before TEM imaging. Further investigating the mechanism of virus transfer at the ultrastructural level would need serial sectioning techniques such as SEM array tomography or block face imaging to reconstruct large volumes, hence increasing the probability to capture rare events within their cellular context.

The composition of ECM is diverse but falls into two classes of macromolecules, namely, proteoglycans and fibrous proteins (Järvelainen et al., 2009; Schaefer and Schaefer, 2010; Stavolone and Lionetti, 2017). Glycosaminoglycan (GAG) chains composed of HS and chondroitin sulfate fill the majority of the ECM (Stavolone and Lionetti, 2017). Several viruses, including herpesviruses, have been shown to interact with HS in a charge-based and a relatively non-specific manner (Azab and Osterrieder, 2017a; Barth et al., 2003; Chen et al., 1997; Tyagi et al., 2001). Herpesvirus envelope glycoproteins (gC and gB) interact reversibly with HS, which lead to virus concentration at the cell surface (Azab and Osterrieder, 2017a) and can explain the embedding of EHV-1 within the ECM. We further showed that virus particles are colocalizing with agrin and collagen of the ECM. The HS proteoglycan, agrin, plays a key role in efficient cell-to-cell communication at the synaptic zone (Alfsen et al., 2005; Dustin and Dustin, 2001; Khan et al., 2001) and in virus cell-to-cell transmission (Alfsen et al., 2005; Dityatev and Schachner, 2006; Pais-Correia et al., 2010). It is worthy to mention that the role of synapse formation in EHV-1 spread has not yet been elucidated. Collagens are the most abundant structural element of the ECM and play a role in the regulation of cell adhesion, providing tensile strength and supporting chemotaxis and migration (Rozario and Desimone, 2010). The actual interaction of the virus with the two proteins and their exact role in virus transmission need to be elucidated. We barely detected a colocalization with ezrin protein, which is a member of the ECM protein family and plays a key role in cell surface structure adhesion, migration, and organization (Vaheri et al., 1997).
The chemical compositions (GAGs and fibrous proteins) of ECM endowed this matrix with its gelatinous properties that play important physiological roles (Rhodes and Simons, 2007). To investigate the functional relevance of ECM integrity on virus spread, we applied mechanical and/or chemical (through heparin or heparinase enzyme) shearing forces to the infected cells. Partial destruction of EHV-1-associated ECM, although did not completely remove virus-ECM structures, significantly reduced virus cell-to-cell spread and the overall infectious capacity of PBMC. Cell supernatants obtained after ECM destruction, on the other hand, contained more virus assemblies and gained more infectivity even in the presence of neutralizing antibodies. Interestingly, destruction of EHV-4-associated ECM did not result in any infectivity at both levels (cell-to-cell or supernatants). Taken together, this indicates that the integrity of ECM is essential for EHV-1 transfer at the PBMC-EC interface. Further studies will be needed to unravel the composition, structure, regulation, and function of the ECM of PBMC. EHV-1 was able to egress the ECM at certain time point, probably when reaching the EC, through a virus-specific mechanism, which is not available for its close relative (EHV-4).

EHV-1 interaction with the ECM components of PBMC and its subsequent egress when reaching the EC is enigmatic. Several herpesvirus glycoproteins have been implicated in cell-to-cell spread (Azab and Osterrieder, 2017a). Two candidates (gB and gD) have been shown to have essential roles in virus entry, penetration, and tissue tropism. In all members of Herpesviridae family, gB is a highly conserved fusogenic protein and it interacts with HS or other specific cell receptors. gD, when present, is the main receptor-binding protein that triggers the entry steps (Azab and Osterrieder, 2017a). Swapping gB and gD between EHV-1 and EHV-4 resulted in a dramatic change in virus infectious ability. ECM-associated EHV-1 with either gD4 or gB4 showed a significant reduction in cell-to-cell spread. On the other hand, EHV-4 with gB1 or gD1 obtained, for the first time, the ability to spread from infected PBMC to EC. We surmise that EHV-1 gD and gB were able to interact with certain structures of the ECM, such as HS or yet unidentified cell receptors, that keep the virus attached during the whole trip. How the virus breaks this interaction to infect the EC is unknown. It is possible that the virus interacts with EC while attaching to the ECM of PBMC. Another explanation could be that the virus adheres avidly to the ECM deposited by EC before penetration. It is possible that following virus binding to the ECM of PBMC, the ECM could be remodeled resulting in the virus having greater affinity for the ECM of EC. Such interaction with EC can be stronger and specific to influence virus transfer from a temporary situation on the surface of PBMC to a stable one on EC. The clear difference between the fusogenic functions of EHV-1 and EHV-4 gBs can also explain the difference of infection outcomes (Spiesschaert et al., 2019b). The interaction of EHV-1 gD with new cell receptors on the surface of EC cannot be excluded. Detection of EHV4 associated with the ECM indicates that the virus can interact or, at least, be captured by the gelatinous nature of the ECM. However, the failure of virus egress to infect EC can be interpreted by the absence of suitable genetic makeup that can support its release. Being stuck longer may lead to its inactivation on the surface of PBMC as a defense mechanism as previously shown with Newcastle disease virus (Yaacov et al., 2012). This interpretation is supported by the ability of EHV-4 to egress and infect EC after supporting its genome with more powerful genes (gD and gB) from EHV-1.

Transcellular and paracellular pathways have been described to illustrate PBMC migration through the endothelial barrier (Engelhardt and Wolburg, 2004; Muller, 2011). Although the process of transcellular migration and the influencing factors are not fully understood, the entire process of trafficking through the endothelium requires three basic steps: rolling, adhesion, and transmigration (Schmidt et al., 2011). We have recently shown that EHV-1-infected PBMC have the ability to roll and then adhere to the endothelium to facilitate virus spread, probably after the upregulation of certain adhesion molecules and cytokines (Holz et al., 2017; Proft et al., 2016; Spiesschaert et al., 2015a). Here, we document the final step of trafficking and show that infected PBMC can migrate through the endothelium in a process that might lead to a safe delivery of the virus inside the cell. Although such a mechanism may not represent the default pathway of virus transmission, the possibility to be an “alternative” pathway cannot be excluded. Transmigration of peripheral mononuclear cells containing bacteria through epithelial cells was previously reported (Wewer et al., 2011). It was shown that infection with Streptococcus suis enhanced the process of transcellular migration. Whether EHV-1 has an influencing effect remains to be investigated. We noticed that the transmigrating infected PBMC was digested within a cytoplasmic vacuole. It seems that lysis of naïve PBMC during transmigration is an expected process (Figure S13) (Bedoya et al., 1969; Herman et al., 2019). However, it is also possible that virus particles on the surface of PBMC may trigger cell lysis. As transendothelial migration is a rare event and cannot be easily captured by TEM, we could not detect intact PBMC with virus particles on the surface; however, we detected virus particle on the surface of
digested PBMC. We surmise that virus particles are able to escape from the vacuole before complete lysis through fusion with the vacuole membrane, a process that is well mastered by herpesviruses (Atanasiu et al., 2010).

Cell-cell fusion is considered the classical and most investigated event of herpesvirus transmission (Cole and Grose, 2003). EHV-1-infected PBMC were capable of transferring infected cytoplasmic material into uninfected EC through fusion events. These results are described before for other herpesviruses including HCMV, PRV, and a VZV (Digel et al., 2006; Gerna et al., 2000; Van de Walle et al., 2003) as well as other viruses (Duprex et al., 1999; Takeuchi et al., 2003). In general, herpesviruses are equipped with a set of glycoproteins (gD, gB, gH, and gL) that carry out all fusion events either between the virus and cells or between infected and non-infected cells (Ambrosini and Enquist, 2015; Atanasiu et al., 2010). During viremia, infected PBMC slow down, roll over the endothelium, and finally adhere to EC. Such a process gives the infected cells a chance to establish cell-cell fusion due to replication of intracellular virus particles and expression of viral envelope glycoproteins on the surface of infected PBMC. Interestingly, all detected fusion events were in PBMC infected for 24 h, where viruses already entered the cells and produced the needed glycoproteins for fusion (Van de Walle et al., 2003).

In conclusion, among all viruses, herpesviruses have proven themselves as “masters of co-evolution.” It is surprising how herpesviruses manipulate infected PBMC with different mechanisms and pathways to ensure a safe and secure spread in the same scene. (1) “PBMC-hitching”: at the first hit, herpesviruses attach to the ECM, hide, transport, and then egress to infect other cells. (2) “Intracellular delivery”: transendothelial migration of infected PBMC may lead to a direct delivery of the virus inside the cytoplasm of EC. (3) “Classical-fusion”: this event is well controlled by herpesviruses and always occurs to facilitate virus spread.

Limitations of the Study
Animal study is the main limitation of this study. It is difficult to determine virus spread from PBMC to EC in vivo. One possible animal study could be the infection of horses with EHV-1RFP and collecting PBMC at different time points during viremia and check whether the virus could be present on the surface of the cells. However, this will not prove that the virus was stuck from the beginning on the surface of PBMC. Another limitation is the confirmation that viruses delivered intracellularly with migrating PBMC are infectious. To conduct such experiment, we will need to isolate EC with infected PBMC inside and propagate them separately to track if they will develop infection; however, this is technically very difficult. Finally, characterization of the ECM of PBMC and determining their composition, structure, regulation, and function would add significantly to our understanding of virus spread. However, conducting this experiment was severely affected by the current COVID-19 pandemic, which hindered our regular access to the laboratories.

Resource Availability
Lead Contact
Further information and requests for resources and reagents or data should be directed to and will be fulfilled by the Lead Contact, Walid Azab: walid.azab@fu-berlin.de.

Materials Availability
This study did not generate new unique reagents.

Data and Code Availability
This study did not generate/analyze datasets/code.

METHODS
All methods can be found in the accompanying Transparent Methods supplemental file.

SUPPLEMENTAL INFORMATION
Supplemental Information can be found online at https://doi.org/10.1016/j.isci.2020.101615.
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AUTHOR CONTRIBUTIONS

First author M.K. designed and conducted most of the experiments, designed the figures, and wrote the manuscript. Coauthor S.P. designed and conducted the live cell imaging and prepared the samples for EM, designed live cell imaging figures and movie, and wrote part of the manuscript. Coauthor B.F. designed and conducted the EM experiment. Coauthor T.M. designed and conducted the EM experiment, conducted data analysis of the EM pictures, and wrote part of the manuscript. Corresponding author W.A. designed the project, discussed the results at different stages of work, and wrote and corrected the manuscript.

DECLARATION OF INTERESTS

The authors declare no competing interests.

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Supplemental Information

Equid Herpesvirus-1 Exploits the Extracellular Matrix of Mononuclear Cells to Ensure Transport to Target Cells

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Supplemental Information

Supplemental figures and legends
Figure S1. Colocalization of virus particles with the carbohydrate-rich extracellular matrix, Related to Figure 1. PBMC were infected with EHV-1\textsuperscript{RFP} (red; MOI=0.5) for 1 hr (a) and 24 hr (b and c). Cell surface glycoproteins of the ECM were stained green with FITC-labeled ConA. PBMC nucleus was stained with DAPI (blue). After 24 hr, EHV\textsuperscript{RFP} viral particles were detected outside the cells colocalizing with ECM (b) as well as inside the PBMC (c, arrowheads). Data are representatives of three independent experiments. Scale bar = 10 μm and scale bar of magnification = 7 μm. Image stacks (number of stacks = 17 with 0.75 μm z-stack step size) were photographed using VisiScope Confocal FRAP microscope. Presented here is a single optical section of the stacks. (d) 3D overview of embedded viruses in the ECM. Red: RFP-labeled virus particles; green: FITC-labeled ConA. (D1 and D2) are the front overviews, (D3 and D4) are the side overviews and (D5 and D6) are the back overviews. The 3D images were reconstituted from total projection of stack series image planes (number of stacks = 17) with 0.75 μm z-stack step size.
Figure S2. Colocalization of virus particles with the carbohydrate-rich extracellular matrix after 7 days of infection, Related to Figure 1. PBMC were infected with EHV-1^{RFP} (red; MOI=0.5) for 24 hr and up to 7 days. At different time points (day 1, 2, 3, 5 and 7) PBMC surface glycoproteins of the ECM were stained green with FITC-labelled ConA. Nucleus was stained with DAPI (blue). Data are representatives of three independent experiments. Scale bar = 30 µm and scale bar of the magnification = 5 µm. Image stacks (number of stacks = 15 with 0.5 µm z-stack step size) were photographed using VisiScope Confocal FRAP microscope. Presented here is a single optical section of the stacks.
Figure S3. Colocalization of virus particles with the carbohydrate-rich extracellular matrix, Related to Figure 1. PBMC were infected with EHV-1RFP (MOI=0.5) for 5 min. Cell surface glycoproteins of the ECM were stained green with FITC-labelled ConA. Plasma membrane was stained with CellVue® Claret Far Red Fluorescent dye (red) and nucleus was stained with DAPI (blue). Two independent representative experiments are shown (a-d and e-f). EHV-1RFP viral particles (yellow) were detected outside cells colocalizing with ECM (c and g), but not with plasma membrane (b and f). Scale bar = 30 µm and scale bar of the magnification = 5 µm. Image stacks (number of stacks = 15 with 0.5 µm z-stack step size) were photographed using VisiScope Confocal FRAP microscope. Presented here is a single optical section of the stacks.
Figure S4. Extracellular matrix components of non-infected PBMC, Related to Figure 1. Cell surface glycoproteins of the non-infected PBMC stained green with FITC-labeled ConA (a), lectin from Triticum vulgaris (WGA; b), anti-collagen (c), anti-agrin (d), or anti-ezrin (e). PBMC nucleus was stained with DAPI (blue). Data are representatives of three independent experiments. Scale bar = 10 μm and scale bar of magnification = 7 μm. Image stacks (number of stacks = 17 with 0.75 μm z-stack step size) were photographed using VisiScope Confocal FRAP microscope. Presented here is a single optical section of the stacks.
Figure S5. Modulation of ECM on the surface of infected and non-infected PBMC, Related to Figure 1. PBMC were either mock-infected or infected with EHV-1<sup>RFP</sup> (MOI=0.5) for 5 min, 1 hr, or 24 hr. The cells were stained green with FITC-labeled ConA (a) or lectin from Triticum vulgaris (WGA; b). The experiment was performed three independent times in blinded fashion. The y-axis indicates the total number of cells involved in the assay. Kruskal–Wallis with Dunns test was used to compare modulation of ECM on infected cells at different time points to the non-infected cells; no significant difference was detected. (c) Confocal images showing non-homogenous (A and B) and homogenous (C and D) distribution of ECM stained with ConA (green). Nucleus was stained with DAPI (blue). Image stacks (number of stacks = 17 with 0.75 μm z-stack step size) were photographed using VisiScope Confocal FRAP microscope. Presented here is a single optical section of the stacks.
Figure S6. Virus transmission from infected PBMC to EC, Related to Figure 2. Number of counted viral particles embedded in the ECM of PBMC and transferred to EC is shown. PBMC were infected with EHV-1RFP (MOI=0.5) for 5 min and added to endothelial cells for 2 hr followed by fixation with PFA 4% and image stacks (number of stacks = 17 with 0.75 μm z-stack step size) were photographed using VisiScope Confocal FRAP microscope. (a) Number of counted viral particles on the surface of PBMC and EC. (b) Number of PBMC and EC on which virus particles were present. The experiment was conducted one time in triplicate.
Figure S7. Live cell imaging of virus transmission from infected PBMC to endothelial cells, Related to Figure 2. Time-lapse live cell imaging confocal microscopy showing EHV-1RFP+GFP viral particles transfer from the overlaid 5-min-infected-PBMC on endothelial cells. Viral particles (red, arrowhead) were visualized moving from infected PBMC to endothelial cells. Virus replication in EC is indicated by GFP expression, which is integrated in the viral genome (green) and production of new progenies RFP-labeled virus (red) in the infected cells. At time points 04:08 and 05:28 min, the green pseudocolor was removed to clearly expose virus-RFP signals. Image stacks (number of stacks = 15 with 0.5 μm z-stack step size) were photographed using VisiScope Confocal FRAP microscope. Presented here is a single optical section of the stacks.
**Figure S8. EHV-1 transfer between PBMC, Related to Figure 2.** PBMC infected with EHV-1\textsuperscript{RFP} (red; MOI=0.5) for 1 hr and stained with ConA (green) display aggregates of two PBMC with the likelihood of virus transfer in between. Data are representatives of three independent experiments. Scale bar = 10 μm and scale bar of magnification = 7 μm. Image stacks (number of stacks = 17 with 0.75 μm z-stack step size) were photographed using VisiScope Confocal FRAP microscope. Presented here is a single optical section of the stacks.
Figure S9. Virus particles on the surface of infected-PBMC, Related to Figure 3. Transmission electron microscopy (TEM) of EC grown on gridded coverslips and overlaid with EHV-1RFP infected-PBMCs. Infected cells were observed either interacting with (a and b) or in close proximity (c and d) to EC. TEM images of the targeted PBMC show virus particles (white arrows) on the surface of infected PBMC. Marked parts with rectangles in (a and c) are magnified in (b and d, respectively). Scale bar = 1 μm.
Figure S10. Transcellular migration of infected-PBMC through endothelial cells, Related to Figure 5. Ultrathin cross sections of PBMC crossing endothelial cells. PBMC were infected for 5 min with EHV-1^RFP and incubated over adherent EC for 3 hr. Partially digested infected-PBMC (at different stacks) is shown (a, d, g) and marked with a rectangle, which is magnified (b-c, e-f, h-i). Virus particles attached to migrating PBMC is marked with white arrows. Scale bar =1 μm.
**Figure S11. Expression of viral envelope glycoproteins (gD and gB) on the surface of infected PBMC, Related to Figure 6.** PBMC were infected with EHV-1<sup>RFP</sup> (red; MOI=0.5) for 24 hr and probed with antibody against EHV-1 gD (a-c) or gB (d-f) followed by anti-mouse Alexa Fluor-488 (green). Nucleus was stained with DAPI (blue). Red signals on cell surface and within the cytoplasm or the nucleus of the infected PBMC indicates RFP-labelled virus particles and expression of viral capsid proteins. Data are representatives of three independent experiments. Scale bar = 30 µm and scale bar of the magnification = 5 µm. Image stacks (number of stacks = 15 with 0.5 µm z-stack step size) were photographed using VisiScope Confocal FRAP microscope. Presented here is a single optical section of the stacks.
Figure S12. Colocalization of EHV-4 with the carbohydrate-rich extracellular matrix, Related to Figure 7. (a) PBMC were infected with EHV-4GFP (MOI=0.5) for 5 min. Cell surface glycoproteins of the ECM were stained with FITC-Labeled ConA (green). EHV-4 particles were stained with anti-glycoprotein D monoclonal antibodies (red). PBMC nucleus was stained with DAPI (blue). Data are representatives of three independent experiments. Scale bar = 10 μm and scale bar of magnification = 7 μm. Image stacks (number of stacks = 17 with 0.75 μm z-stack step size) were photographed using VisiScope Confocal FRAP microscope. Presented here is a single optical section of the stacks. (b and c) EHV-4 transmission from infected PBMC to EC under “static” and “dynamic” states. PBMC were infected with EHV-4GFP for 5 min. Infected PBMC were added to EC under “static” conditions (b) or allowed to flow over EC “dynamic” (c) in the presence of neutralizing antibodies. After 24 hr, virus spread was assessed by counting the plaques on EC. As a control, infected PBMC were placed into a transwell insert without direct contact between EC and PBMC “no contact”. The data represent the mean ± standard deviation of three independent and blinded experiments. (d) PBMC were infected with EHV-4GFP (MOI=0.1) for 5 min. The ECM was disrupted with both mechanical (extensive pipetting) and chemical (heparin treatment) together (EHV-4_5M_Cell_pip_hep). The disrupted cells (d.1) or the supernatant (EHV-4_5M_Sup_pip_hep; d.2) were added to EC. As a control, the ECM was left undisrupted (EHV-4_5M_Cell or EHV-4_5M_Sup). As a control of pipetting and heparin treatment, EHV-1 was used. The data represent the mean ± standard deviation of three independent and blinded experiments with EHV-4. Cells were infected with EHV-1 once to confirm virus spread and infection (see Fig 4a and b). (e) Endothelial cells are permissive for EHV-4 cell-free viruses. EC were infected with EHV-4GFP (MOI 0.1) for 1 hr. Virus and media were removed and the cells were overlaid with methylcellulose media 1.5% for 3 days. EHV-4 GFP-expressing viral plaques were imaged with Carl Zeiss AXIO imager microscope equipped with a Zeiss Axiocam. Scale bar = 70 μm.
Figure S13. Transcellular migration of PBMC through endothelial cells, Related to Figure 5.

Ultrathin cross section of non-infected PBMC crossing endothelial cells. Partially digested naïve PBMC is shown within a cytoplasmic vacuole of EC. The migrating PBMC marked with a rectangle in (a, red arrow) is magnified in (b, red arrow). Scale bar = 1 μm.
Transparent Methods

Methods

Viruses. All viruses used in this study were produced and recovered from infectious bacterial artificial chromosome (BAC) clones. Those were BACs of EHV-1 (EHV-1^{GFP}) strain Ab4 (Goodman et al., 2007), EHV-4 (EHV-4^{GFP}) strain TH20p (Azab et al., 2009), recombinant EHV-1_gB4 and EHV-4_gB (Spiesschaert et al., 2015b), and EHV-1_gD4 and EHV-4_gD1 (Azab and Osterrieder, 2012). Viruses were grown on equine dermal (ED) cells (CCLV-1222, Federal Research Institute for Animal health, Germany) (Spiesschaert et al., 2015b). All recombinant viruses express the enhanced green fluorescent protein (EGFP; encoded in the mini-F sequence under the control of the immediate-early CMV promoter) for efficient identification of infected cells. All virus stocks were prepared by infecting ED cells until having 100 % cytopathic effect (CPE). Infected cultures (supernatant and cells) were collected and centrifuged at 6000 x g for 5 min to get rid of cellular debris. Virus-rich supernatants were collected and stored at -80 °C.

mRFP1-labeled viruses. Insertion of monomeric red fluorescent protein (mRFP1) into VP26 of EHV-1 Ab4 strain was performed as described before (Azab et al., 2013b; Tischer et al., 2006). Briefly, mRFP1 was amplified by PCR using pEPmRFP1-in (Tischer et al., 2006) as a template; all used primers are referenced in (Azab et al., 2013b). The resulting PCR products were electroporated into GS1783 (a kind gift from Dr. Greg Smith, Northwestern University, Chicago, IL) harboring the corresponding BACs (EHV-1 with and without GFP; to give rise of EHV-1^{RFP, GFP} and EHV-1^{RFP}, respectively). Kanamycin-resistant colonies were purified and screened by PCR and restriction fragment length polymorphism (RFLP) analyses using EcoRV and BamHI. Positive clones were subjected to a second round of Red recombination to obtain the final constructs after excision of kanamycin gene. The final clones were confirmed by PCR, RFLP and sequencing (data not shown). All viruses (EHV-1^{RFP, GFP} and EHV-1^{RFP}) were reconstituted as described before (Azab et al., 2013b).

Cells. Primary equine carotid artery endothelial cells (a kind gift from Prof. Dr. Johanna Plendl, Freie Universität Berlin, Institut für Veterinär-Anatomie) were propagated in Dulbecco’s modified Eagle’s medium (DMEM; PAN-Biotec Ltd. Germany) supplemented with 20% fetal bovine serum (FBS) (PAN-Biotec Ltd. Germany), 1% nonessential amino acids (Biochrom, Germany), and 1% penicillin-streptomycin. Endothelial cells were obtained according to the rules of Institutional Animal Care and Committee of Berlin (Landesamt für Gesundheit und Soziales, L 0089/14). Equine PBMC were isolated from heparinized blood collected from healthy horses by density gradient centrifugation over Biocoll L6715 (Biochrom, Germany), following the manufacturer’s instructions. After two washing steps, cells were resuspended in RPMI-1640 (Pan Biotech, Germany) supplemented with 10% FBS, 0.3 mg/ml glucose, nonessential amino acids, and 1% penicillin-streptomycin. Blood collection was performed according to the rules of Institutional Animal Care and Committee of Berlin (Landesamt für Gesundheit und Soziales, L 0294/13). PBMC were negative for both EHV-1 and EHV-4 as tested by qPCR (data not shown). PBMC were either used fresh after isolation or preserved in liquid-nitrogen and reconstituted when needed.

Plasmids. For assessing fusion assay by luciferase activity measurement, we used pT7EMCLuc plasmid, which expresses firefly luciferase reporter gene under the control of T7 promoter, and pCAGT7 plasmid, which expresses T7 RNA polymerase. Both plasmids were kindly provided by Dr. Richard Longnecker, Northwestern University. A plasmid expressing mRFP under the control of Human CMV promoter, pEP-CMVmRFP-in (Campbell et al., 2002; Tischer et al., 2006), was used for fusion assay. pEPmRFP1-in carrying kanamycin mRFP cassette had been used as a shuttle plasmid for EHV-1 strain Ab4-mRFP-VP26 construction.

Antibodies. FITC-labeled ConA (C7642), Triticum vulgaris (WGA; L4895) were from Sigma. Mouse anti-agrin antibody (D-2: sc-374117) and mouse anti-ezrin antibody (3C12: sc-58758) were from Santa Cruz Biotechnology. Rabbit anti-human Collagen I/II/III/IV/V was from (Biorad: 2150-2206). Mouse anti-EHV-4 gD Mab was kindly provided by Jules Minke, Merial: 105E2K (Azab et al., 2013a). Mouse Anti-EHV-1 gD 19-mer polyclonal antibodies (Allen and Yeargan, 1987) were kindly provided by Dr. Dennis O’Callaghan, Louisiana State University Health Sciences Center, Shreveport, LA. Mouse EHV-1 anti-gB monoclonal antibodies (mAb) (Wellington et al., 1996) were kindly provided by Dr. Udeni B. R. Balasuriya, Gluck equine research center, University of Kentucky, KY.
Confocal microscopy and immunofluorescence. For detection of the extracellular carbohydrate-rich matrix, 2x10^6 PBMC (in suspension) were infected with EHV-1^RFP^ or EHV-4^GFP^ (MOI=0.5) for 5 min, 1 hr, 24 hr, 2 days, 3 days, 5 days, and/or 7 days. Cells were centrifuged at 400 x g for 5 min and treated with ice cold citrate buffer (pH 3) for 1.5 min. The cells were washed three times with PBS and fixed with 4% Parafomaldehyde (PFA) for 20 min. For EHV-4, virus particles were stained with mouse anti-EHV-4 gD monoclonal antibody followed by Alexa Fluor 568-labeled goat anti-mouse IgG (1:1000; Invitrogen; 1841756). The carbohydrate-rich extracellular matrix was stained with FITC-labelled ConA and WGA (dilutions were 1:250 and 1:1000, respectively). Collagen, agrin and ezrin were stained by rabbit anti-human Collagen I/III/IV/V, mouse anti-agrin antibody, and mouse anti-ezrin antibody with dilutions of 1:100, 1:50, and 1:50, respectively, followed by Alexa Fluor 488-labeled goat anti-rabbit (1:1000; Invitrogen; 1966932) and anti-mouse IgG (1:1000; Invitrogen; 2015565). Plasma membrane was stained with CellVue® Claret Far Red Fluorescent dye (Sigma-Aldrich). DAPI (Life Technologies) was used to stain the nucleus. Image stacks (number of stacks = 15-17 with 0.5-0.75 µm z-stack step size) were photographed using VisiScope Confocal FRAP microscope (<40 magnification) and equipped with ANDOR iXon Ultra 888 camera (Visitron Systems GmbH, Germany). The images were processed and analyzed by using Image J software (https://imagej.nih.gov/ij/) and Metamorph software 7.8 (Molecular Devices).

In another experiment, 1x10^6 PBMC were infected with EHV-1^RFP^ (MOI=0.5). All infected cells were treated with ice-cold citrate buffer and virus-neutralizing antibodies as described above. The cells were added to endothelial cell monolayers for 2 hr. The cells were fixed and stained with FITC-labeled ConA and WGA. Image acquisition, processing and analysis were performed as mentioned above.

Flow chamber assay. The assay was performed as previously described (Spiesschaert et al., 2015b). Briefly, PBMC (1x10^6) were infected with EHV-1^GFP^ or EHV-4^GFP^ (MOI=0.5) for different time points (5 min, 1 hr, 6 hr). PBMC were treated with ice-cold citrate buffer (pH 3). The infected cells were washed two times and resuspended in medium containing 1:100 dilution of virus neutralizing antibody (titer: 1:2028 as determined by serum neutralization test (Azab et al., 2019) at 37°C for 30 min (Goehring et al., 2011; Spiesschaert et al., 2015a). Endothelial cells were grown to confluency in 0.4 collagen IV-coated cell flow chambers (Ibidi Integrated BioDiagnostics) that were connected to a perfusion system by Luer locks (Ibidi Integrated BioDiagnostics). The flow chamber set was incubated at either 4°C or 37°C. Infected PBMC were introduced at a flow rate of 0.5 mm/s, which is within the mammalian physiological range (0.34 to 3.15 mm/s) (Hudetz et al., 1996). The velocity was calculated according to the size of the chamber and the velocity in mammalian brain capillaries and was generated by a NE-4000 double-syringe pump (New Era Pump System). After 24 hr, GFP-viral plaques on the EC monolayer were counted (excluding the inlet/outlet areas) using an inverted fluorescence microscope and equipped with AxioCam MRc camera (Zeiss Axiosvert 100).

To evaluate virus transfer from different PBMC subpopulations (i.e. T lymphocyte, B lymphocyte and monocytes) to EC, each PBMC subpopulation was sorted out and flow chamber assay was performed. Briefly, PBMC were stained with (1:200 diluted) primary mouse antibodies against equine CD3 (T lymphocyte), IgM (B lymphocyte) and CD14 (monocyte). The antibodies were kindly provided by Dr. Bettina Wagner, Cornell University, Ithaca, NY, USA. PBMC were then labelled with secondary Alexa Fluor 488-conjugated goat anti-mouse IgG antibody (Invitrogen) and sorted using FACSaria (BD Bioscience™). Sorted T- lymphocytes, B-lymphocytes and monocytes (1x10^6) were infected with EHV-1^GFP^ (MOI=0.5) for 5 min at 37 °C. After incubation, cells were treated with citrate buffer, resuspended in EHV-1 neutralizing antibodies and flow chamber experiment was performed as described above.

Contact assay. PBMC (5x10^5) were infected with EHV-1^GFP^ or EHV-4^GFP^ at MOI of 0.1 at different time points (5min, 1 hr, and 6 hr). Infected cells were treated with ice-cold citrate buffer and added to EC for 2 hr in the presence of virus neutralizing antibody as described above. Non-adherent cells were then removed by extensive and gentle washing with PBS. In the "no-contact" setup, virus-infected PBMC were placed into a transwell insert (with a membrane pore size of 0.4 µm; Corning Transwell support system) without direct contact with EC. Methylcellulose 1.5% was added and cells were incubated for 24 hr. Virus transfer was assessed by counting the green fluorescent viral plaques using an inverted fluorescence microscope (Zeiss Axiosvert 100).
**Disruption of extracellular virus assemblies.** PBMC (5x10^5) were infected with EHV-1\textsuperscript{RFP+GFP} or EHV-4\textsuperscript{GFP} at MOI of 0.1 for 5 min as described above. The cells were subjected to vigorous pipetting, treated with heparin, or both. Heparin (50μg/ml; Sigma-Aldrich) was added to the cells for 30 min. PBMC (cells or supernatant) were added to EC for 2 hr. After washing, methylcellulose 1.5% was added to the cells. Viral plaques on EC were counted after 24 hr using an inverted fluorescence microscope. For confocal microscopy, EHV-1-infected PBMC were stained with FITC-labeled ConA and then treated with heparinase III (10 U ml\textsuperscript{-1}) for 1 hr or heparin (50μg/ml) for 30 min at 37°C in serum-free medium. Image stacks were photographed using VisiScope Confocal FRAP microscope. Image acquisition, processing and analysis were performed as mentioned above.

**Fusion assays.** Luciferase assay. Day 1: PBMC were nucleofected with 2μg pCAGT7 plasmid using Amaxa® Cell Line Nucleofector® Kit V (Lonza) according to the manufacturer instructions. Day 2: PBMC were infected with EHV-1\textsuperscript{GFP} at MOI of 1. Endothelial cells were nucleofected with 2μg pT7EMCLuc and seeded in collagen-coated 24 well plate. Day 3: Nucleofected-infected-PBMC were washed and overlaid on the nucleofected endothelial cells. Day 4: Activation of the reporter luciferase gene resulted from fusion of nucleofected infected PBMC to nucleofected endothelial cells was measured by Luciferase Assay System kit (E1500, Promega, USA) using TriStar LB 941 Multimodal plate reader (Berthold Technologies) for luminescence measurement according to the manufacturer's instructions (Promega). Briefly, endothelial cells were washed to remove non-fused PBMC and lysed with the lysis buffer. The lysed cells were left on a shaker for 15 min and 20 μl of the cell lysates were used for the reading. As negative controls, only nucleofected PBMC or only infected PBMC were overlaid on the nucleofected endothelial. As a positive control, PBMC were nucleofected with both plasmids (pCAGT7 and pT7EMCLuc).

In another assay, PBMC were nucleofected with 2 μg pEP-CMVmRFP-in as described above. After 24 hr, nucleofected PBMC were infected with EHV-1\textsuperscript{GFP} at MOI of 1. After 24 hr, nucleofected-infected PBMC were added to endothelial cells for 3.5 hr. Endothelial cells were washed and fixed with 4% PFA and analyzed using VisiScope Confocal FRAP microscope (×40 magnification). Images were pseudocolored according to their respective emission wavelengths, overlaid and processed using Image J and Metamorph softwares.

To determine the expression of fusogenic viral glycoproteins (gB and gD) on the surface of infected PBMC, cells were infected with EHV-1\textsuperscript{RFP} (MOI=0.5) for 24 hr. Infected cells were fixed with 4% PFA for 20 min and stained with primary mouse antibody against EHV-1 gB and gD (dilution of 1:250) for 1 hr. Goat anti-mouse Alexa Fluor-488 was used as secondary antibody and nucleus was stained with DAPI. Cells were imaged using VisiScope Confocal FRAP microscope as described above.

**Live cell imaging.** PBMC were infected with EHV-1\textsuperscript{RFP+GFP} (MOI=3) for 5 min then the cells were treated with ice-cold citrate buffer and virus neutralizing antibody. The infected cells were added to endothelial cells grown on 8-wells ibidi slide coated with collagen (Ibidi Integrated BioDiagnostics). Cells were imaged in non-phenol red DMEM cell culture media (Pan Biotech, Germany) supplemented with 20% FBS, 1% penicillin and streptomycin, and 1% non-essential amino acid. The temperature on the microscope stage was held stable during time-lapse sessions using an electronic temperature-controlled airstream incubator. The field area was chosen displaying EHV-1\textsuperscript{RFP+GFP} viral particles (red signals) on the surface of PBMC overlaying the EC. Images were captured in time-lapse every 8 min time series using a VisiScope Confocal FRAP microscope (×20 magnification). Images and movies were generated and analyzed using ImageJ and Metamorph softwares. Long time-lapse experiments were carried out using the autofocus function integrated into the advanced time series macro set.

**Correlative fluorescent and electron microscopy.** For TEM analysis, endothelial cells were grown on correlative microscopy gridded coverslips (LCMC34A, Plano GmbH, Germany). PBMC were infected with EHV-1\textsuperscript{RFP} (MOI=5) for 5 min then the cells were treated with ice-cold citrate buffer and virus neutralizing antibody. The infected-PBMC were overlaid on EC and incubated at 37°C for 1 or 3 hr. Cells were fixed with 4% PFA for 30 min at room temperature. Cell nuclei were counter-stained with 2μg/ml Hoechst 33342 (Thermo Fisher Scientific, Germany) in PBS for 10 min at room temperature and kept in PBS after staining. Cells covering the region of one grid square on the coverslip were imaged using a CellDiscoverer 7 microscope (Carl Zeiss, Germany) at 20x magnification. For TEM, cells were then fixed on the coverslips in 2.5% glutaraldehyde (Grade I, Sigma, Germany) freshly prepared in PBS from a 25%
stock solution (overnight at 4°C). After washing 3 times in PBS, coverslips were incubated in 0.5 % (v/v) osmium tetroxide (Science Services GmbH, Germany) in PBS for 1 hr at room temperature, washed 4 times for 20 min in distilled water, incubated for 30 min in 100 mM Hepes buffer containing 0.1 % (w/v) tannic acid (Science Services, Germany), washed 3 times for 10 min in distilled water and then again incubated in 2 % (w/v) uranyl acetate (Sigma-Aldrich, Merck, Germany) for 1.5 hr at room temperature. After washing once in distilled water, samples were then dehydrated through a series of increasing ethanol concentrations (5 min in 30 %, 10 min in 50 %, 15 min each in 70%, 90%, 96% and finally 3 times 10 min in absolute ethanol, respectively). Samples were then incubated overnight at 4°C in 100 % SPURR resin (Low Viscosity Spurr Kit, Ted Pella, CA, USA). After exchanging fresh SPURR resin 2 times (2 hr at room temperature), coverslips were mounted between two pieces of Aclar Embedding Film (Science Services, Germany) for polymerization (24 hr at 60°C). For TEM imaging, areas (comprising approximately 3x3 squares from the grid pattern) exhibiting 2-4 PBMC carrying a virus load closely attached to EC were cut out, mounted on a preformed resin dummy and embedded again in SPURR resin in order to cut target cells with an orientation perpendicular to coverslip support. After polymerization, sample blocks were carefully trimmed close to the selected cells using a EM Trim2 specimen trimmer (Leica Microsystems, Germany) followed by cutting 70 nm sections using a UC7 ultramicrotome (Leica Microsystems, Germany) equipped with a 3 mm diamond knife (Diatome, Biel, Switzerland). Sections were placed on 3.05 mm carbon-coated Formvar copper slot grids (Plano GmbH, Germany) and post-contrasted using 2% (w/v) uranyl acetate and Reynolds lead citrate. TEM images were taken at 1100x - 6500x nominal magnification using a Tecnai Spirit transmission electron microscope (FEI) operated at 120 kV and equipped with a 4k x 4k F416 CMOS camera (Tietz Video and Image Processing Systems GmbH; TVIPS).

**Statistical analyses.** Statistics were performed using SPSS 23 and GraphPad Prism 7. Normally distributed data sets, determined by the Shapiro-Wilk test, were analyzed with one-way analysis of variance (ANOVA) followed by Dunnett's test for multiple comparisons. Data sets that were not normally distributed were analyzed with Kruskal-Wallis test followed by Dunns test for multiple comparison tests.

*, $P < 0.05$, **, $P < 0.01$, ***, $P < 0.001$. 
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