Mycobacteria employ two different mechanisms to cross the blood–brain barrier

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Abstract
Central nervous system (CNS) infection by Mycobacterium tuberculosis is one of the most devastating complications of tuberculosis, in particular in early childhood. In order to induce CNS infection, M. tuberculosis needs to cross specialised barriers protecting the brain. How M. tuberculosis crosses the blood–brain barrier (BBB) and enters the CNS is not well understood. Here, we use transparent zebrafish larvae and the closely related pathogen Mycobacterium marinum to answer this question. We show that in the early stages of development, mycobacteria rapidly infect brain tissue, either as free mycobacteria or within circulating macrophages. After the formation of a functionally intact BBB, the infiltration of brain tissue by infected macrophages is delayed, but not blocked, suggesting that crossing the BBB via phagocytic cells is one of the mechanisms used by mycobacteria to invade the CNS. Interestingly, depletion of phagocytic cells did not prevent M. marinum from infecting the brain tissue, indicating that free mycobacteria can independently cause brain infection. Detailed analysis showed that mycobacteria are able to cause vasculitis by extracellular outgrowth in the smaller blood vessels and by infecting endothelial cells. Importantly, we could show that this second mechanism is an active process that depends on an intact ESX-1 secretion system, which extends the role of ESX-1 secretion beyond the macrophage infection cycle.

KEYWORDS
blood–brain barrier, ESX-1 secretion, Trojan horse mechanism, tuberculosis, tuberculous meningitis, zebrafish

1 | INTRODUCTION

Tuberculous meningitis (TBM) is one of the most severe extra-pulmonary manifestations of tuberculosis (TB) and significantly contributes to mycobacterial disease burden (WHO, 2017). Invasion of Mycobacterium tuberculosis, the causative agent of TB, into the central nervous system (CNS) occurs in 1% of all cases (Thwaites, van Toorn, & Schoeman, 2013; Wilkinson et al., 2017). Major risk groups for developing TBM include young children and HIV-positive individuals in TB endemic areas (van Well et al., 2009; Wilkinson et al., 2017). Despite
extensive research efforts, the diagnosis and treatment of TBM is often delayed because of its insidious onset (Wilkinson et al., 2017). Consequently, half of the patients are diagnosed in the most advanced stage of disease, resulting in a high mortality rate of nearly 20% and neurological sequelae in more than half of the survivors (Chiang et al., 2014). These poor odds of (full) recovery for TBM patients can be mostly attributed to the severe neuro-inflammation at the base of the brain, on-going neural ischaemia, and vasculitis (Donald & Van Toorn, 2016).

The histological hallmark of TB is the granuloma, a cluster of immune cells that shields off the infected macrophages from the surrounding tissue. In 1933, it was established that in TBM, granulomas are present in brain parenchyma and meninges. This important observation led to the hypothesis that granulomas were the main aetiology of TBM and these infectious foci were later called Rich foci (Rich & McCordock, 1933; Rich & Thomas, 1946). Today, the concept of the Rich focus still stands; however, the question remains how the first mycobacterium enters the brain to seed the Rich focus.

To induce granuloma formation and subsequently meningitis, \( M. \text{tuberculosis} \) must traverse the blood–brain barrier (BBB), a selectively permeable layer that separates brain tissue from the blood circulation. The BBB consists of specialised endothelial cells connected by tight junctions, closely surrounded and monitored by several cell types, including astrocytes, pericytes, and microglia. The BBB regulates the passage of molecules and effectively protects the brain from circulating toxins and microorganisms (Abbott, Patabendige, Dolman, Yusof, & Begley, 2010; Abbott, Rönnbäck, & Hansson, 2006; Obermeier, Daneman, & Ransohoff, 2013). Little is known about the steps preceding granuloma formation, in particular how \( M. \text{tuberculosis} \) manages to traverse the BBB.

Only a small subset of bacterial pathogens is able to cause meningitis or CNS infections. Thus far, three different BBB traversal strategies have been described for these pathogens. The most commonly utilised route is transcellular migration. This receptor-mediated process results in endocytosis of the pathogen by endothelial cells that line the blood vessels and is used by \( \text{Streptococcus pneumoniae} \), \( \text{Haemophilus influenzae} \), and \( \text{Neisseria meningitidis} \) (Bencurova, Mlynarcik, & Bhide, 2011; Kim, 2008; Orihuela et al., 2009; van Sorge & Doran, 2013). A second route is paracellular migration, which usually occurs when BBB integrity is disrupted by direct contact with the pathogen or as a result of secreted bacterial toxins. A third mechanism of BBB crossing is the Trojan horse mechanism; the pathogen infects a macrophage that subsequently traverses the BBB. Based on the fact that \( M. \text{tuberculosis} \) is an intracellular pathogen capable of surviving and replicating within the macrophage, the latter mechanism seems logical for BBB traversal (Nguyen & Pieters, 2005). In line with this hypothesis, \( M. \text{tuberculosis} \) was found to cross an epithelial barrier with significantly higher efficiency when phagocytosed by monocytes than when mycobacteria alone were introduced in an in vitro system (Bermudez, Sangari, Kolonoski, Petrofsky, & Goodman, 2002). Furthermore, macrophages played an essential role in early dissemination and establishment of extra-pulmonary foci (Clay et al., 2011; Polena et al., 2016). However, the ability of \( M. \text{tuberculosis} \) to invade brain endothelial cells in vitro has been described as well (Be, Bishai, & Jain, 2012; Chen, Sakamoto, Quinn, Chen, & Fu, 2015; Jain, Paul-Satyaseela, Lamichhane, Kim, & Bishai, 2006; Mehta, Karls, White, Ades, & Quinn, 2006). Consequently, the exact mechanisms involved in mycobacterial invasion into brain tissue are still not completely understood. We reasoned that, in order to study such a detailed sequence of events, it is essential to observe the interplay between host and pathogen in vivo.

Several in vivo models to study mycobacterial pathogenesis exist, such as rabbits, guinea pig, and mice (Be et al., 2012; Be, Klinkenberg, Bishai, Karakousis, & Jain, 2011; Skerry et al., 2013; Tsenova et al., 2005; Tucker et al., 2016; van Well et al., 2007; Zucchi et al., 2012). However, none of these models could demonstrate the course of events during mycobacterial trafficking across the BBB in a living host. Another in vivo model that has proven to mimic human mycobacterial disease well is the zebrafish—\( \text{Mycobacterium marinum} \) infection model (Berg & Ramakrishnan, 2012; Lesley & Ramakrishnan, 2011; van der Sar, Appelmelk, Vandebroucke-Gruals, & Bitter, 2004). The translucency of the Danio rerio larvae in combination with many fluorescent tools offers unique possibilities to study host–pathogen interaction in real life (Kuipers, Kalicharan, Wolters, van Ham, & Giepmans, 2016; Tobin, May, & Wheeler, 2012). Moreover, the anatomy of the zebrafish BBB is highly similar to the human BBB. Already after 3-days postfertilisation (dpf), the zebrafish BBB functionally prevents exchange of large molecules (Fleming, Diekmann, & Goldsmith, 2013; Xie, Farage, Sugimoto, & Anand-Apte, 2010). Importantly, upon infection with \( M. \text{marinum} \), TBM does occur in adult zebrafish, with granuloma formation in the meninges and brain parenchyma in 20% of the cases (van Leeuwen et al., 2014). Therefore, the zebrafish model allows us to specifically address questions regarding mycobacterial invasion into the brain in an in vivo model (Bemut et al., 2014; Tenor, Oehlerls, Yang, Tobin, & Perfect, 2015; van Leeuwen et al., 2014).

This study provides in vivo evidence that mycobacteria utilise phagocytic cells to cross the BBB. Additionally, by using in vivo macrophage depletion and correlated light and electron microscopy (CLEM), we show that mycobacteria also employ transcellular migration by infecting and damaging brain endothelial cells in an ESX-1-dependent manner.

2 | RESULTS

2.1 | \( M. \text{marinum} \) crosses a functionally intact BBB within phagocytic cells

To examine the importance of an intact BBB in mycobacterial trafficking to the brain, we used larvae at different developmental stages and followed infection progression daily (Figure 1a). BBB biogenesis in zebrafish starts at 3 dpf and can be determined by systemic injection of fluorescent tracers. A 3-kDa fluorescent dye was excluded from the larval brain from 3 dpf onwards, indicating BBB maturation (compare Figure 1b with 1c). Please notice that the blood vessels in close proximity to the eyes and gills do not possess a BBB and therefore do not restrict diffusion of the dye into the surrounding tissue (Figure 1c; van Leeuwen et al., 2017).

Infection experiments performed at 2 dpf, that is, before BBB biogenesis, showed that mycobacteria readily crossed blood vessel walls in the brain at this time point (Figure 1d,e). In these larvae, mycobacterial migration was observed as early as 1-day postinfection (dpi; Figure 1d). Examination of larvae infected at 4 dpf, that is, after the formation of
the BBB, showed that mycobacteria are present in brain blood vessels at 1 and 2 dpi (corresponds to 7 dpf), infected with *M. marinum* E11:mEos3.1 (green). Red arrow marks the caudal vein injection spot. Boxed area represents the brain region of which representative images are shown in this figure. (b) *Tg(kdrl:mCherry)y5* zebrafish larva, uninfected, 1 hr after 3-kDa Alexa 680 tracer injection at 2 dpf, showing massive leakage of tracer to the ventricles, confirming the immaturity of the BBB. (c) Injection of tracer at 7 dpf, showing localisation in brain blood vessels, indicating that the BBB is functionally intact at this moment. Blood vessels in close proximity to the eyes and gills do not possess a BBB and do not restrict diffusion of the dye, leading to extravasation of the dye at these locations. Scale bars bc = 250 μm. (d) *Tg(kdrl:mCherry)y5* zebrafish larvae infected at 2 dpf (before formation of a functional BBB), 1 dpi single nonphagocytosed bacteria (green) can be found in blood vessel (red; open arrow) and in brain tissue (arrow), showing that *M. marinum* can enter brain tissue at this time point. (e) Also at 2 dpi, phagocytosed bacteria (colocalisation of green bacteria and cyan phagocytes labelled with anti-L-plastin) in blood vessel (open arrow) and single nonphagocytosed bacteria in brain tissue (arrow) are found. (f) Systemic infection at 4 dpf (larvae with a functional BBB), *M. marinum* (green) is solely found in brain blood vessels (red) at 1 dpi and (g) 2 dpi (arrows). (h–j) From 3 dpi onwards, phagocytosed mycobacteria (colocalisation of green bacteria and cyan phagocytes) are able to leave the bloodstream (arrows). (k) Representative image of section of brain in *Tg(Fli1:GFP)y1* zebrafish larvae, systemically infected with the esx-1 mutant (red) at 4 dpf, showing that mutant bacteria also cross the BBB within phagocytic cells (anti-L-plastin, cyan). (lm) Examples of spot with high expression of mCherry tagged VEGFr2 in *Tg(kdrl:mCherry)y5* larva, colocalising with phagocytosed (cyan) *M. marinum* E11 (green) in blood vessel (red). (l) Merged image, (m) corresponding single red channels. Scale bars d–m = 25 μm

**FIGURE 1** *M. marinum* WT and esx-1 mutant traverse across an intact blood–brain barrier within macrophages. (a) Lateral view of a casper zebrafish larva at 3 dpi (corresponds to 7 dpf), infected with *M. marinum* E11:mEos3.1 (green). Red arrow marks the caudal vein injection spot. Boxed area represents the brain region of which representative images are shown in this figure. (b) *Tg(kdrl:mCherry)y5* zebrafish larva, uninfected, 1 hr after 3-kDa Alexa 680 tracer injection at 2 dpf, showing massive leakage of tracer to the ventricles, confirming the immaturity of the BBB. (c) Injection of tracer at 7 dpf, showing localisation in brain blood vessels, indicating that the BBB is functionally intact at this moment. Blood vessels in close proximity to the eyes and gills do not possess a BBB and do not restrict diffusion of the dye, leading to extravasation of the dye at these locations. Scale bars bc = 250 μm. (d) *Tg(kdrl:mCherry)y5* zebrafish larvae infected at 2 dpf (before formation of a functional BBB), 1 dpi single nonphagocytosed bacteria (green) can be found in blood vessel (red; open arrow) and in brain tissue (arrow), showing that *M. marinum* can enter brain tissue at this time point. (e) Also at 2 dpi, phagocytosed bacteria (colocalisation of green bacteria and cyan phagocytes labelled with anti-L-plastin) in blood vessel (open arrow) and single nonphagocytosed bacteria in brain tissue (arrow) are found. (f) Systemic infection at 4 dpf (larvae with a functional BBB), *M. marinum* (green) is solely found in brain blood vessels (red) at 1 dpi and (g) 2 dpi (arrows). (h–j) From 3 dpi onwards, phagocytosed mycobacteria (colocalisation of green bacteria and cyan phagocytes) are able to leave the bloodstream (arrows). (k) Representative image of section of brain in *Tg(Fli1:GFP)y1* zebrafish larvae, systemically infected with the esx-1 mutant (red) at 4 dpf, showing that mutant bacteria also cross the BBB within phagocytic cells (anti-L-plastin, cyan). (lm) Examples of spot with high expression of mCherry tagged VEGFr2 in *Tg(kdrl:mCherry)y5* larva, colocalising with phagocytosed (cyan) *M. marinum* E11 (green) in blood vessel (red). (l) Merged image, (m) corresponding single red channels. Scale bars d–m = 25 μm.

Because BBB crossing was only observed after 3 days after intravenous infection, we hypothesised that possibly at this stage, the BBB might be compromised due to the infection and a concomitant inflammatory response. In order to study the integrity of the BBB during initial mycobacterial migration to brain tissue, we injected a 3-kDa fluorescent dye at 3 and 5 dpi and monitored dye distribution 60–180 min postinjection. At all time points, we found many single bacteria associated with the blood vessel wall, possibly in the process of migration, with fluorescent dye restricted to the vessel lumen (Figure S2A,B,D open arrow). The lack of leakage of dye in the parenchyma suggests an intact BBB at these spots. However, injection of dye at 3 and 5 dpi resulted in accumulation and colocalisation of dye within these clusters (Figure S2A,C,E,F closed arrow). This indicates that once an inflammatory focus is formed, the local integrity of the BBB is...
reduced. Because we did not observe accumulation of dye in the ventricles, we reasoned that there is no increased overall leakage and no substantial breakdown of the BBB in this inflammatory setting.

To further study the Trojan horse as migration mechanism, we compared *M. marinum* WT with a mutant strain (ecCcB1:tn) deficient in ESX-1 secretion (esx-1 mutant). ESX-1 secretion mutants are severely attenuated (Davis & Ramakrishnan, 2009; Stoop et al., 2011; Volkman et al., 2004), but most importantly for our purposes, these mutants are unable to complete the macrophage infection cycle and are therefore predominantly located inside phagocytic cells (Houben et al., 2012; Simeone et al., 2012; van der Wel et al., 2007).

As expected, we observed significantly lower numbers of mycobacteria in zebrafish larvae infected with our esx-1 mutant (compare Figure 1j [WT: 28 single infected phagocytes and 14 early clusters in six larvae] with 1k [esx-1 mutant: 16 single infected phagocytes and three small clusters in eight larvae]; see also Figure 5a,b). The observed bacteria were always associated with phagocytes (Figure 1k). Despite these lower numbers, esx-1 mutants were still able to infect brain tissue and in both WT and esx-1 mutant infected larvae, approximately half of the infected macrophages were found in brain parenchyma (WT: 13/28, esx-1 mutant: 8/16). Collectively, our findings confirm the protective function of the BBB against *M. marinum* infection of brain tissue in developing zebrafish larvae and indicate that *M. marinum* also uses phagocytes to cross the BBB. In addition, we have indications that local BBB integrity seems to be reduced once an inflammatory focus is established.

### 2.2 | Intensified VEGFR2 signal colocalises with infected phagocytes

Previously, it has been shown that upregulation of vascular endothelial growth factor (VEGF) had a promoting effect on macrophage-medi-ated extra pulmonary dissemination of *M. tuberculosis* (Polena et al., 2016). To examine the role of VEGF in our zebrafish model, we systemically infected *Tg(kdr:mCherry)JS* embryos with *M. marinum*. These embryos are modified to express mCherry under control of the promoter regulating *kdr/ vegfr2* gene expression, which allowed us to determine the effect of *M. marinum* on *vegfr2* expression in blood vessels of the brain. We observed that, from 3 dpi, the VEGFR2 signal was more intense at 54% of the spots in which phagocytosed mycobacteria colocalised with blood vessels (Figure 1l,m, 13 out of 24 spots in 20 larvae). This time course corresponds with the observed migration of mycobacteria into brain tissue. Nonphagocytosed bacteria found in brain blood vessels showed significantly lower colocalisation with an intensified VEGFR2 signal (16% of the cases, data not shown). This observation is indicative for local upregulation of vegfr2 in endothelial cells by phagocytes carrying mycobacteria.

We hypothesised that alterations in general VEGF levels could affect the percentage of migrated infected phagocytes. Therefore, we manipulated VEGF levels using two different approaches, (a) inducing VEGF levels with GS4012 (Wu et al., 2015) or (b) blocking the VEGF receptor with SU5416 (Oehlers et al., 2014; Wu et al., 2015). Although differences in overall infection levels were found with an increase in the absolute number of infected cells after inducing VEGFR2 signalling (Figure S3), the proportion of infected macrophages crossing the BBB at 3 dpi was similar for all groups (GS4012: 48%; control: 52%; SU5416: 52%; Table S1).

Taken together, although we do observe a local upregulation of VEGFR2 at the site of BBB crossing by infected phagocytes, generic manipulation of the VEGF levels does not alter the percentage of migrated phagocytosed mycobacteria in zebrafish embryos.

### 2.3 | Wild type *M. marinum* can still infect brain tissue when phagocytes are depleted

To examine whether the Trojan horse mechanism forms the only transport route to cross the BBB, we studied mycobacterial invasion in parenchyma of zebrafish larvae that were depleted of phagocytes. Successful depletion was achieved by injecting both pu.1 morpholinos at the single cell stage, to prevent macrophage development (Clay et al., 2011) and clodronate-filled liposomes at 3 dpf to kill the remaining circulating phagocytic cells (Figure 2a,b; Bernut et al., 2014; Pagan et al., 2015; van Rooijen, Sanders, & van den Berg, 1996).

As expected, infection with wild type *M. marinum* in control larvae with normal phagocyte counts resulted in clusters of infected macrophages in the brain of zebrafish larvae (Figure 2c–f). In these zebrafish, we even identified mycobacteria-loaded phagocytes that appear to be in the process of crossing the BBB (Figure 2d,f, arrow). In contrast, infection in larvae without phagocytes resulted in a huge expansion of mycobacteria in blood vessels without the formation of early granulomas in brain tissue (Figure 2g–j). Surprisingly, however, mycobacteria were also still present in brain tissue in all cases (Figure 2h,i, indicated with *). This observation suggests that mycobacteria can utilise another, phagocyte-independent, route to cross the BBB. Closer analysis of the bacterial aggregates showed colocalisation with Fli1 labelling, which labels endothelial cells (Lawson & Weinstein, 2002; Figure 2h,j, arrow), suggesting mycobacterial outgrowth in other cell types than phagocytes. In addition, the normal vascular architecture seemed to be disrupted at these heavily infected spots, which indicates major changes in endothelial cells. Although mycobacteria were still located in brain tissue in these larvae, we observed a completely different pattern and distribution of infection (compare Figure 2k with 2l). In phagocyte-depleted larvae, *M. marinum* was always found in close vicinity of blood vessels in the brain that were highly loaded with bacteria, whereas in untreated zebrafish, granulomas were located at more distant locations indicating that phagocytes are essential for transport of bacteria into deeper tissue.

In conclusion, *M. marinum* has the capability to migrate into brain tissue in the absence of phagocytes, which means that an alternative migration route is present.

### 2.4 | Wild type *M. marinum* causes damage to blood vessels and surrounding tissue

To be able to understand the phagocyte-independent interaction with the BBB in more detail, we used CLEM, which facilitates the direct correlation of fluorescent confocal microscopy with electron microscopy of consecutive slides of the same tissue (Figures 3a–c and 4a–c).

In the wild type situation, *M. marinum* is found to cross the BBB and invade brain tissue, apparently without disrupting the integrity
**FIGURE 2** Wild type *M. marinum* can still infect brain tissue when macrophages are depleted. Left panel shows untreated zebrafish larvae (control), whereas right panel shows larvae depleted of phagocytes by treatment with pu.1 and clodronate filled liposomes at 3 dpf, to kill circulating phagocytes. (a) Control casper larva at 5 dpf, stained with anti-L-plastin to visualise normal phagocyte distribution. (b) Five dpf phagocyte-depleted casper larva. Anti-L-plastin is used in (a) and (b) to stain phagocytes. (c) Dorsal view of wild type *Tg(Fli1:GFP)y1* larvae (green) at 5 dpi, systemically infected at 4 dpf with *M. marinum* (red) and stained with anti-L-plastin (cyan), showing formation of early granuloma in brain tissue. (d) Z-stack of boxed area in (c), allowing for a more precise analysis of the position of *M. marinum* and phagocytes. (e) Corresponding red fluorescent channel to clearly show infection pattern. (f) In the presence of macrophages, *M. marinum* leaves the bloodstream within phagocytes (arrow) and forms early granulomas in brain tissue. Scale bar c = 100 μm. Scale bar d-f = 25 μm. (g) Z-stack of head of *Tg(Fli1:GFP)y1* larva, phagocyte-depleted, and systemically infected with *M. marinum* (red) at 4 dpf and stained with L-plastin (cyan). Boxed area is enlarged in h-j. (h) Shows mycobacteria outside blood vessels in brain tissue in the absence of phagocytic cells (*). (i) Shows the corresponding red fluorescent channel, depicting that tissue infiltration follows the shape of the blood vessels. (j) Single Z-slice, which shows the intracellular phenotype of *M. marinum* (arrow), colocalising with a blood vessel, but in the absence of L-plastin labelling. Scale bar g = 100 μm; Scale bar h-j = 25 μm. (k) Schematic representation of pooled data of all early granulomas found in nine wild type larvae showing a random distribution in the brain. (l) Schematic representation of pooled data of infection distribution found in nine phagocyte-depleted larvae, showing that mycobacteria are found in brain tissue, but do not migrate into deeper tissue.
FIGURE 3  Correlative light and electron microscopy of *M. marinum* infected blood vessels showing irregular blood vessel walls and invasion of endothelial cells. Tg(kdrl:mCherry)155 larva (9 dpf) with red fluorescent blood vessels infected with green fluorescent *M. marinum* after phagocyte depletion. To aid correlation of confocal and EM imaging, nuclei were stained with DAPI after fixation (cyan). (a) Electron microscopy and (b) correlative light and electron microscopy, and (c) confocal microscopy. Arrows indicate landmarks used to merge images obtained from consecutive slices in the same area of the zebrafish brain. Boxed area is enlarged in (f), scale bar a = 5 μm (applies to b–d). (d) Single red channel illustrating the irregularity of the infected blood vessel and the more regular shape of the non-infected blood vessel (right upper corner). (e,f) High magnification EM image showing the irregular shaped infected blood vessels and basal lamina. Red dotted lines represent basal lamina found in this area. Boxed area is enlarged in (g). Scale bar = 1 μm. (g) High magnification of area where mycobacteria can both be found intravascular as intracellular. Vesicles, indicative for intravascular localisation, can only be found left of the red dotted line. This suggest that mycobacteria right of the line are localised in an endothelial cell (*). Scale bar = 200 nm. M = *M. marinum*
of the blood vessel. Bacteria-loaded phagocytes are clearly detected outside of the intact vessels (Figure S4A–C). In contrast, the phenotype found in phagocyte-depleted larvae is completely different. Notably, individual infected blood vessels were shaped irregularly (Figure 3d–f), whereas non-infected blood vessels appear intact (Figure 3d, upper-right corner). Furthermore, several blood vessels seemed to be segmented, visible within a cross section of a vessel (Figure 3d–f). Higher magnifications of these infected blood vessels showed that bacteria can be found both intravascular and intracellular. For example, in Figure 3g, the bacteria are located in an endothelial cell (specified with *), as the red dotted line indicates the membrane separating an endothelial cell from the vascular lumen. Furthermore, we observed that infection with wild type *M. marinum* disrupts the vessel wall (Figure 4c,f) and consequently the basal lamina (Figure 4f,g).

Taken together, we show that, in the absence of phagocytes, mycobacteria are capable of invading surrounding tissue, presumably the endothelial cells, and induce damage to the basal lamina and local distribution in the surrounding brain tissue.

### 2.5 ESX-1 secretion is essential for macrophage-independent BBB crossing

Next, we examined the fate of our esx-1 mutant in macrophage-depleted zebrafish larvae. In line with previous findings (Clay et al., 2011), we observed significantly higher outgrowth of the esx-1 mutant under these conditions, as compared with outgrowth in normal zebrafish larvae (Figure 5a,b), confirming that the absence of phagocytes compensates for the attenuation of this strain. Closer examination of phagocyte-depleted larvae revealed an important difference with WT *M. marinum* infection: Although blood vessels in the brain were filled and clogged with ESX-1-deficient mycobacteria (Figure 5c,d), esx-1 mutants were only rarely found outside the blood vessels. (Figure 5e,f, 16/61 cases in five larvae). Therefore, esx-1 mutants seem unable to cross the BBB efficiently under these conditions, with subsequent bacterial outgrowth in the vessel lumen and protrusion of vessels (Figure 5g,h, open arrow). Only occasionally, we observed bursting of blood vessels (Figure 5g,h, closed arrow). Therefore, for macrophage-independent crossing, ESX-1 secretion seems to be an important factor.

Also for the esx-1 mutant infections, CLEM analysis was performed in phagocyte-depleted larvae (Figure 6a–c), which confirmed that high amounts of extracellular bacteria were present in brain blood vessels. The esx-1 mutant mycobacteria were predominantly found in the lumen of the blood vessels and were never found to cross or disrupt the basal lamina and blood vessel wall (Figure 6d, red dotted line), which is in contrast with WT infections. Only sporadically, bacteria were endocytosed by an endothelial cell (Figure 6e,*). In addition, no segmentation of blood vessels was observed, although the
infected vessels were often enlarged in diameter (Figure 6f; lumen diameter WT infection: average 6.2 μm, range 4–9 μm, n = 15; lumen diameter esx-1 mutant infection: average 18.4 μm, range 7.5–37 μm, n = 15).

Collectively, CLEM analysis confirmed that mycobacteria require ESX-1 secretion for macrophage-independent crossing of the BBB.

2.6 | *M. marinum* invasion of brain endothelial cells is dependent on ESX-1 secretion

To study the interaction of *M. marinum* with brain endothelial cells (BECs) in more detail and to examine the role of ESX-1 secretion in this process, human BECs were infected with live or heat-killed
M. marinum WT, the esx-1 mutant or the complemented esx-1 mutant. Bacterial levels at 24 hr postinfection (hpi) were analysed with FACS and compared with uptake of these bacteria by RAW macrophages. RAW macrophages phagocytosed both strains with similar efficiency, both at 2 and 24 hpi (Figure 7a), confirming previous reports that the level of phagocytosis of M. marinum WT and our esx-1 mutant are similar (Houben et al., 2012; Simeone et al., 2012; van der Wel et al., 2007).

When compared with macrophages, infection of BECs with M. marinum WT is less efficient. To observe infection, we needed a higher multiplicity of infection (MOI 50 instead of MOI 10) and even then the number of infected cells was lower compared with the RAW macrophages (compare 45% infected RAW macrophages in Figure 7a with 30% infected BECs in Figure 7b). However, if we just look at BECs, we could observe significant differences between M. marinum WT and the esx-1 mutant (Figure 7b). The esx-1 mutant showed almost eight times lower uptake at 24 hpi, for both experiments performed with an MOI of 10 and an MOI of 50. Importantly, the infection defect was restored when cells were infected with the complemented strain. Interestingly, infection with heat-killed M. marinum WT also resulted in reduced uptake, at a level comparable with the phenotype seen for the esx-1 mutant. This indicates that invasion of BECs by M. marinum WT is an active process that depends on an active ESX-1 secretion system.

Confocal analysis of infected BECs showed colocalisation of WT M. marinum with LAMP1, a late endosomal marker, demonstrating that M. marinum E11 is transported to the phagolysosome (Figure 7c,d). In contrast, the few esx-1 mutant bacteria that were associated with

![Image](99x336 to 496x735)
BECs did not colocalise with LAMP1 and based on the 3D modelling appeared to be located at the cell periphery, probably even at the cell surface (Figure 7e,f).

In conclusion, these experiments show that *M. marinum* is capable of actively invading and infecting brain endothelial cells. In addition, the ESX-1 secretion system is essential for active invasion of BECs,
extending the role of ESX-1 secretion beyond the macrophage infection cycle.

3 | DISCUSSION

In this study, we show that *M. marinum* employs two different strategies to cross the BBB: the Trojan horse mechanism and ESX-1-dependent invasion and damaging of endothelial cells (Figure 8, graphical abstract).

Our observation that, under normal conditions, all bacteria that cross the BBB colocalise with a phagocyte is a strong indication that this is one of the mechanisms for *M. marinum*. There was a clear difference in *M. marinum* trafficking before and after the presence of an intact BBB. Infection experiments at early time points, that is, performed before BBB formation, showed that mycobacteria could readily traverse into brain tissue. These results are comparable with mycobacterial invasion into tissue outside the CNS (Lesley & Ramakrishnan, 2011). After the formation of the BBB, mycobacterial crossing was delayed by several days. This shows that the BBB is an obstacle for entering the brain, even when using the Trojan horse mechanism. This also means that in vivo experiments performed to study dissemination of mycobacteria at early time points, that is, 2 dpf, cannot be used to study natural infection of brain tissue.

For extra-cellular pathogens causing meningitis, such as *S. pneumoniae*, *N. meningitides*, and *H. influenza*, it has been shown that haematogenous dissemination directly results in acute meningitis (Panackal, Williamson, van de Beek, Boulware, & Williamson, 2016; van Sorge & Doran, 2013). However, it seems likely that the initial seeding of *M. tuberculosis* into brain tissue only induces a mild inflammatory response, because massive inflammation seen during clinical presentation of TBM is only observed when a pre-existing granuloma in the brain progresses into meningitis (Donald, Schaaf, & Schoeman, 2005; Donald & Van Toorn, 2016; Panackal et al., 2016). This might be explained by the Trojan horse mechanism but also by a sporadic bacteraemia with subsequent non-inflammatory deposition of free mycobacteria on the endothelium. Using a macrophage as protective niche instead of endothelial cell infection, *M. tuberculosis* avoids interaction with the endothelial cells and a subsequent inflammatory response. Our fluorescent dye experiments support that this initial non-inflammatory event is not accompanied with substantial breakdown of the BBB. However, we cannot exclude the possibility of local, perhaps transient, areas of compromised BBB integrity, serving as entry point for infected phagocytes. Additional evidence for a Trojan horse mechanism was obtained by studying esx-1-deficient bacteria. Infection with this attenuated mutant strain that is not able to escape the phagocyte resulted in reduced infection levels and inflammation, but nevertheless, phagocytes filled with these bacteria did infect the brain. Extensive breakdown of the BBB by a massive (local) inflammatory response is unlikely in this case, because we have relatively low levels of infection by this attenuated mutant. Both concepts fit with the generally accepted Rich focus theory about TBM pathogenesis, describing that meningitis only occurs when a granuloma in brain tissue or meninges discharges its content in the subarachnoid space. However, more recently, it has been debated that haematogenous dissemination in the form of miliary TB in young children, with a high bacteraemia, has a high likelihood to lead to TBM soon after infection and may have a stronger correlation with the onset of TBM in children than initially thought (Donald et al., 2005). Therefore, it has been suggested that *M. tuberculosis* might be able to use other entry routes into the brain when high bacterial levels in blood are present. Moreover, mycobacteria associated with endothelium and in close proximity of blood vessels could explain endovasculitis as major pathological hallmark of TBM seen in historical studies and might suggest a specific role of these bacteria in Rich focus formation (Rich & McCordock, 1933).

Our new data uncover the presence of such an alternative route, where *M. marinum* is able to enter the CNS without the help of phagocytes. We used CLEM to show the localisation of mycobacteria in the blood vessel lumen, in endothelial cells, and in the surrounding tissue. These experiments suggested that these bacteria cross the BBB transcellular. Our in vivo studies are consistent with in vitro studies demonstrating the ability of *M. tuberculosis* to invade endothelial cells (Chen et al., 2015; Jain et al., 2006; Mehta et al., 2006). Remarkably, this transcellular migration by *M. marinum* seems to differ from the “classical” transcellular migration route, in which changes in endothelial barrier integrity are generally not observed (Kim, 2008). However, strategies that affect BBB integrity to gain direct entry have been demonstrated for other pathogens (van Sorge & Doran, 2013), and it is not unlikely that mycobacteria developed similar strategies. The
perception that *M. tuberculosis* is also invading other host cells than only the phagocytic cells is not new. Recently, lymphatic endothelial cells were suggested as a replicative niche for *M. tuberculosis* (Lerner et al., 2016), and also the importance of the interaction of *M. tuberculosis* with epithelial cells for early dissemination has been noted previously (Menozzi et al., 2006). Here, we show the role of differential host cells for dissemination across the BBB.

We used an ESX-1-deficient mutant to study the effect of a well-known virulence factor on BBB trafficking. Interestingly, while our results confirmed the importance of ESX-1 secretion in dissemination and virulence (Stoop et al., 2011; Volkmann et al., 2004), it also revealed a completely new and different phenotype. In phagocyte-depleted larvae infected with the ESX-1-deficient strain, a high bacterial load is found in the zebrafish brain. However, in contrast to WT bacteria, these bacteria are almost exclusively restricted to the blood vessel lumen and do not invade endothelial cells. Previously, the work of Jain et al. showed a difference between *Mycobacterium bovis* BCG and *M. tuberculosis* isolates in invasion efficiency of brain endothelial cells, but they did not attribute this difference to ESX-1 (Jain et al., 2006). Our novel findings now show that invasion and infection of endothelial cells by *M. marinum*, both in vivo and in vitro, are dependent on ESX-1 secretion. Our data suggest that ESX-1 substrates mediate an active process in which *M. marinum* is able to invade endothelial cells. This attributes a novel function to ESX-1 secretion, in addition to phagosomal escape (Abdallah et al., 2011; Houben et al., 2012; Simeone et al., 2012; van der Wel et al., 2007). Furthermore, the damaging effect of mycobacteria on endothelial cells might be part of the explanation for the extensive vasculopathy, including stroke and vasculitis, found in clinical TBM presentation and autopsy material (Donald & Van Toorn, 2016; van der Flier et al., 2004; Zaharie et al., unpublished).

Another factor linked to cerebral vasculopathy is VEGF. In the case of TBM, increased VEGF levels have been found in CSF and are associated with cerebral oedema formation, hydrocephalus, and basal meningeal enhancement (van der Flier et al., 2004; Visser et al., 2014). The observed colocalisation of intensified kdr/vegf2 signal with *M. marinum*-loaded phagocytes in larval brain blood vessels in our study suggests an interaction between infected macrophages and the blood vessel wall and a role for VEGF in this process. It has been shown that macrophages secrete VEGF after infection with *M. tuberculosis* in an ESX-1-dependent manner, which in turn interacts with VEGFR2 specifically (Polena et al., 2016). This mechanism is essential for angiogenic vessel growth and bacterial expansion and dissemination. Our data suggest a similar mechanism involved in BBB crossing. The intensified VEGFR2 signal phenotype is less frequently seen when extracellular bacteria colocalise with blood vessels, suggesting that infected macrophages play an important role in this process.

Although it is clear that VEGF is not the only factor involved in mycobacterial CNS infection, the importance of VEGF is of great interest in the context of clinical TBM presentation and therapeutic approaches. Various animal studies suggest that anti-VEGF treatment, such as the widely prescribed anti-VEGF antibody bevacizumab, can resolve vascular leakiness and improve outcome of TB (Datta et al., 2015; Oehler et al., 2014; Oehler et al., 2016). With the rising awareness that host-directed therapy could be a valuable addition to antibiotics, VEGF seems to be an interesting target. However, our study failed to show any effect of VEGF level manipulation on phagocyte migration. The absence of a link might be obscured by many factors, such as VEGF interacting with VEGFR1 in addition to VEGFR2 or TNFα interacting with VEGFR2 (Jeltsch, Leppänen, Saharinen, & Alltalo, 2013; Zielonka et al., 2011). Hypothetically, these pathways interact with each other and may alter the effect seen in our experiments. This implies that anti-VEGF treatment might not be as straightforward as hoped for or this might suggest that VEGF levels are not essential for mycobacterial CNS invasion.

In summary, our results support the longstanding idea that mycobacteria employ macrophages as Trojan horse as migration mechanism across the BBB and suggest that VEGF might play a role in this process, but we also show that the Trojan horse is not the only route this pathogen uses. We demonstrate an additional type of migration whereby virulent *M. marinum* actively infect and disrupt the endothelium to gain excess to brain tissue in an ESX-1-dependent manner. The next step would be to determine if the same mechanisms apply for *M. tuberculosis*. As such, this study reveals early steps of TBM pathogenesis and might help us to explain variation in pathological and clinical presentation.

4 | EXPERIMENTAL PROCEDURES

4.1 | Bacterial strains and growth conditions

*Mycobacterium marinum* E11 WT, a sea bass isolate of *M. marinum*, and the ESX-1 secretion mutant eccCb1::tn, derivative of this strain (esx-1 mutant), were used in this study (Stoop et al., 2011). *M. marinum* E11 and eccCb1::tn were transformed by electroporation with pSMT3-hsp60-mCherry, to express mCherry, or pSMT3-hsp60-mEos3.1 and pSMT3-hsp60-mEos3.2, to express mEos3.1 or mEos3.2 respectively. All three plasmids were used to visualise infection in zebrafish larvae and human Cerebral Microvascular Endothelium Cells (hCMEC/D3) infection experiments. Complementation of eccCb1::tn was done by introduction of plasmid pMV-hsp60-ecCcM1. Transformants were selected on plates with the appropriate antibiotic selection markers (25-μg/ml kanamycin [Sigma] and/or 50-μg/ml hygromycin [Roche]). All constructs were confirmed by sequencing of plasmid inserts.

Strains were routinely grown at 30 °C on Middlebrook 7H10 agar plates (Difco) supplemented with 10% oleic acid-albumin-dextrose-catalase (BD Bioscience) or in Middlebrook 7H9 broth (Difco) with 10% Middlebrook albumin-dextrose-catalase (BD Bioscience), 0.05% Tween-80, and the appropriate antibiotic selection marker.

4.2 | Construction of plasmids

To visualise bacterial infection in zebrafish embryo, infection experiments pSMT3-hsp60-mEos3.1 and pSMT3-hsp60-mEos3.2 were created. Both mEos3.1 and mEos3.2 were obtained from Zhang et al. (2012). mEos3.1 and mEos3.2 were amplified by PCR with 15 base pair flanking regions corresponding to the target vector using mEos3-FW and mEos3-RV (Table S2). The target vector pSMT3-
The plasmids were subsequently transformed by electroporation into digested vector by introduced a 15 base pair overlap with the target vector on both 5' and 3' sides of the insert to allow In-Fusion cloning. The target vector, pMV-sp60-mEos3.1 (Van De Weerd et al., 2016) was digested with NheI and Clal. Subsequently, the insert was introduced into the digested vector by In-Fusion cloning to produce pMV-hsp60-eccCb1. The plasmids were subsequently transformed by electroporation into M. marinum E11 or the respective M. marinum E11 eccCb1::tn mutant.

### 4.3 Injection stocks

Injection stocks were prepared by growing bacteria until the logarithmic phase (OD\textsubscript{600} of 0.7–1). Bacteria were briefly spun down and washed with 0.3% Tween-80 in phosphate buffered saline (PBS) for declumping and resuspended in PBS with 20% glycerol and stored at −80 °C. Before use, bacteria were resuspended in PBS containing 0.17% (v/v) phenol red (Sigma) to aid visualisation of the injection process.

### 4.4 Zebrafish

Zebrafish lines used in this study: transparent casper zebrafish (White et al., 2008), Tg(Fli1:GFP); casper zebrafish, with green fluorescent endothelial cells (Lawson & Weinstein, 2002), and Tg(kdrt;mCherry); with red fluorescent endothelial cells (Jin, Beis, Mitchell, Chen, & Stainier, 2005).

All procedures involving D. rerio (zebrafish) larvae were performed in compliance with local animal welfare laws and maintained according to standard protocols (zfins.org). The breeding of adult fish and infection of embryos older than 5–7 dpf was approved by the institutional animal welfare committee (Animal Experimental licensing Committee, DEC). All protocols adhered to the international guidelines specified by the EU Animal Protection Directive 86/609/EEC, which allows zebrafish embryos to be used up to the moment of free-living (approximately 5–7 dpf). Infection of older embryos was approved under DEC number MM10-02 by the institutional animal welfare committee (Animal Experimental licensing Committee, DEC) of the VU University medical centre.

### 4.5 Infection procedure

At 2 or 4 dpf, embryos were infected with M. marinum E11 or eccCb1::tn by microinjection in the caudal vein (E11: 200–400 CFU; E11-4dpf: 250–900; eccCb1::tn: 4dpf: 325–1,040 CFU). Injection was performed as described previously (Benard et al., 2012). At 1–5 dpi bacterial infection was monitored daily with a Leica MZ16FA fluorescence microscope. Following analysis, larvae were fixated on indicated time points overnight in 4% (v/v) paraformaldehyde (EMA, 100122) in PBS, and stored in 100% methanol at −20 °C for immune-histochemical staining and confocal imaging. To determine the exact number of bacteria injected, the injection volume was plated on 7H10 plates containing the proper antibiotic selection. During injection and microscopic examining, larvae were anaesthetised in egg water with 0.02% (W/V) buffered 3-amino benzoic acid (Tricaine; Sigma-Aldrich, A-5040).

### 4.6 Phagocyte depletion

To deplete the phagocytic pool in larvae, pu.1 morpholino (Rhodes et al., 2005) was injected at the 1–4 cellular stage. At 3 dpf clodronate-filled liposomes, (1:1 [v/v] diluted in phenol red, 10 nl end volume) were injected into the caudal vein, to deplete the larvae of all residual systemic phagocytes (Bernut et al., 2014; Pagán et al., 2015; Van Rooijen & Sanders, 1994).

### 4.7 BBB functionality assay with fluorescent tracer

Uninfected Tg(kdrt;mCherry);5 larvae were injected with a 3-kDa fluorescent dye (Dextran, Alexa Fluor 647, Thermofisher) into the caudal vein at 2–9 dpf. Leakage of tracer into brain tissue, as a measure for BBB integrity, was subsequently monitored on a confocal microscope every 10 min between 30- and 120-min postinjection.

### 4.8 VEGF modulation experiments

To study the effect of VEGF on infection levels and bacterial BBB crossing, either 250 nM SU5416 (VEGF receptor blocker, Sigma S8442; Oehlers et al., 2014; Wu et al., 2015) or 2.5 μM GS4012 (VEGF inducer, Santa Cruz Biotech sc-222411; Wu et al., 2015) was used. Compounds were added directly after infection and refreshed every second day as described previously (Oehlers et al., 2014). Six to eight larvae per group were fixated in 4% (v/v) paraformaldehyde (EMS, 100122) in PBS at 3 and 5 dpi for further analysis with confocal microscopy.

### 4.9 Immunohistochemical stain

Larvae were fixated on indicated time points overnight in 4% (v/v) paraformaldehyde (EMS, 100122) in PBS and stored in 100% methanol at −20 °C for immune-histochemical staining and confocal imaging. Larvae were labelled with anti-L-plastin, which stains phagocytic cells according to established protocols (Bennett et al., 2001; Herbomel, Thisse, & Thisse, 1999). In short, larvae were rinsed with 1% PBTx, (1% Triton X-100 in PBS), permeated in 0.24% trypsin in PBS and blocked for 3 hr in block buffer (10% normal goat serum [NGS] in 1% PBTx). Samples were incubated with anti-L-plastin (1:500 [v/v] dilution) in antibody buffer (PBTx containing 1% [v/v] NGS and 1% [w/v] BSA) overnight at RT. Samples were washed with PBTx, incubated for 1 hr in block buffer and stained with an Alexa-Flour-647 goat-anti-rabbit as secondary antibody (Invitrogen A21070, 1:400), overnight at 4 °C.

### 4.10 Microscopy

Bacterial infection was monitored initially with a Leica MZ16FA fluorescence microscope. Bright field and fluorescence images were
generated with a Leica DFC420C camera. Early granuloma formation was analysed visually and quantified with custom-made pixel-counting software (www.elaborant.com). Confocal analysis was performed on hCMEC/D3 cells and larvae, embedded in 1% low melting-point agarose (Boehringer Mannheim, 12841221-01) in an eight-well microscopy μ-slide (ibidi). Analysis was performed with a confocal laser-scanning microscope (Leica TCS SP8 X Confocal Microscope). Leica Application Suite X software and ImageJ software were used to adjust brightness and contrast to create overlay images and 3D models.

4.11 CLEM

Representative embryos were selected to study infection in the zebrafish brain using CLEM. Larvae were fixed overnight in 4% (v/v) paraformaldehyde (EMS, 100122) dissolved in PBS, and stored in 0.1-M PHEM and 0.5% PFA. 0.4-M PHEM was made with 240-mM PIPES, dissolved in 0.3-M NaOH. Following this, 40-mM EGTA, 100-mM HEPES, and 8-mM MgCl₂ were added in this order. pH was adjusted to 6.9 with NaOH, and PHEM buffer was stored at ~20 °C until use. Larvae were incubated overnight in 2-M sucrose and snap-frozen on a pin in liquid nitrogen. Semithin and ultrathin EM sections were cut as described by Bedussi et al. (2016). Semithin sections (300–400 nm) were stained with DAPI for counterstaining of nuclei in the tissue and analysed with confocal microscopy. When fluorescent vessels were found to colocalise with fluorescent M. marinum, 70- to 100-nm thin sections were cut and transferred to a 150-mesh copper grid and stained with Uranyl Acetate for TEM analysis, grids with ultrathin sections were washed and stained with a Uranyl acetate/Tylose mixture and imaged using Tecnai T12 at 120 kV. The position of the nuclei, which is visible in both FM and EM, was used to align and overlay the images (Adobe Photoshop CS6).

4.12 Cell infection

The human Cerebral Microvascular Endothelial Cells line hCMEC/D3 (or BECs; Weksler et al., 2005), kindly provided by Dr. P.-O. Couraud (Institut Cochin, Universite Paris Descartes, Paris, France), and mouse RAW264.7 macrophages (American Type Culture Collection) were used for cell infection experiments. The hCMEC/D3 cells were grown in EBM–2 medium supplemented with hEGF, hydrocortison, GA-1000, fetal bovine serum, VEGF, hFGF-B, R-3/IGF-1, ascorbic acid, and 2.5% fetal calf serum (Lonza, Basel, Switzerland). Before use, cells were washed with PBS and human endothelial SFM was added (Invitrogen, ca. no 11111-044). RAW264.7 cells were cultured in RPMI 1640 with Glutamax-1 (Gibco) supplemented with 10% FBS (Gibco), 100 U of penicillin/ml, 100 μg of streptomycin/ml at 37 °C, 5% CO₂.

4.13 Cell infection–flow cytometry

hCMEC/D3 cells were seeded until confluent in 24-well plates. For RAW macrophages, a total of 3 × 10⁷ cells were seeded in T175 flasks (Corning). M. marinum E11, M. marinum eccCb1::tn, and M. marinum eccCb1::tn-comp were grown until the exponential growth phase, spun down, and resuspended in specialised medium. Brain endothelial cells were infected with a multiplicity of infection of 10 or 50, incubated for 2 or 24 hr at 30 °C, 5% CO₂, washed with PBS and detached with trypsin. RAW macrophages were infected with an MOI of 5, 10, or 50 for 2 or 24 hr and incubated at 30 °C, 5% CO₂. Uptake of both strains was quantified for both cell lines with a BD Accuri C6 flow cytometer (BD Biosciences) with a 488-nm laser and 585/-40-nm filter to detect mEos3.1. A minimum of 10,000 gated events were collected per sample per time point; data were analysed using BD CFlow software.

4.14 Cell infection–confocal microscopy

hCMEC/D3 cells were seeded until confluent in eight-well microscopy μ-slide (ibidi, cat no. 80826). M. marinum E11 and M. marinum eccCb1::tn were grown until the exponential growth phase, spun down, and resuspended in specialised medium. Brain endothelial cells were infected with an MOI of 10 or 50, incubated for 2 or 24 hr, washed with PBS and fixed with 4% PFA dissolved in PBS for 20 min. For labelling, cells were blocked for 60 min in block buffer (5% NGS in 0.3% Triton X100). Samples were incubated with anti-LAMP1 (Cell Signalling, cat. no 9091P, 1:100) in antibody buffer (1% BSA and 0.3% Triton X100 in PBS) overnight at 4 °C. Samples were washed with PBS and incubated with Alexa-Fluor-488 goat-anti-rabbit (Molecular Probes, cat. no A-11008, 1:400) in antibody buffer for 90 min at RT. Cells were washed with PBS, incubated with Hoechst (1:1000, Molecular Probes, cat. no 33258) for 1 min, washed with PBS, and stored in PBS at 4 °C until further analysis with confocal microscopy (Leica TCS SP8 X Confocal Microscope). Leica Application Suite X software was used for 3D analysis.

4.15 Graphs and statistical analysis

Graphs were made using Graph Pad Prism 6.0. Pixel counts obtained with eLaborant were logarithmic transformed; error bars represent mean and standard error of the mean. A one-way ANOVA was performed followed by a Bonferroni multiple comparison test to analyse statistical significance. Graphs with results for D3 and RAW cell infection show percentage infected cells of total cells; error bars represent mean and standard error of the mean. A one-way ANOVA was performed for statistical analysis.

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**SUPPORTING INFORMATION**

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