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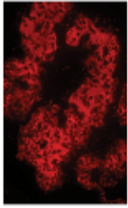
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Targeting of the central nervous system by *Listeria monocytogenes*

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Abbreviations: *Lm*, *Listeria monocytogenes*; CNS, central nervous system; CSF, cerebrospinal fluid; CT, computed tomography; BBB, blood-brain barrier

Among bacteria that reach the central nervous system (CNS), *Listeria monocytogenes* (*Lm*) is one of the deadliest, in human and ruminant. This facultative intracellular bacterium has the particularity to induce meningitis, meningoencephalitis and rhombencephalitis. Mechanisms by which *Lm* accesses the CNS remain poorly understood, but two major routes of infection have been proposed, based on clinical, in vitro and in vivo observations. A retrograde neural route is likely to occur in ruminants upon crossing of the oral epithelium, and this probably accounts for the observation that *Lm* induces almost exclusively rhombencephalitis in these animals. In contrast, the hematogenous route is likely the most frequent in human, in whom bacteria circulating in the blood, either free or associated with leukocytes are thought to breach the blood-brain barrier. New animal models that faithfully reproduce the hallmarks of human neuroinfection will allow addressing the molecular mechanisms underlying *Lm* ability to induce CNS disease, and improve our understanding of the pathophysiology of this deadly infection.

Introduction

Central nervous system (CNS) invasion by bacteria leads to severe infection, which can be fatal or associated with severe sequelae.^{1,2} Among the bacteria that can access the CNS, *Listeria monocytogenes* (*Lm*), a Gram-positive facultative intracellular bacterium, has the ability to induce meningitis and encephalitis.

The first known isolate of *Lm* was obtained from the cerebrospinal fluid (CSF) of a soldier who died from meningitis in 1918. The report was published in 1921,³ before the characterization of a new bacterial species in 1926 from infected laboratory rabbits and guinea pigs⁴ and in 1927 from wild gerbils.⁵ This species is now known as *Listeria monocytogenes*. However, it is only during the second half of the 20th century that listeriosis started to be considered as a prominent human infection, concomitant with changes in food habits and the introduction of immunosuppressive therapies, in addition to being a well-characterized zoonosis routinely observed in domestic ruminants. The first human

listeriosis outbreak linked to the absorption of contaminated food was published in 1983,⁶ and *Lm* is now regarded as a classic foodborne pathogen for veterinarians and clinicians, which is well known to infect ruminants and humans via the oral route, and target similar organs: the central nervous system and the fetoplacental unit.⁷

Even if listeriosis is rare in human (0.1 to 10 cases/million; 0.1% of all foodborne infections), it is considered as the most severe bacterial foodborne infection, responsible for 1,645 cases in Europe in 2009 (4 cases/million) for a case fatality rate of 16.6%⁸ and around 2,500 cases per year in the USA, associated with a lethality of up to 30% in case of neurological involvement, even when appropriately treated.^{9,10} More than 50% of the cases correspond to septicemia, around 20–25% to CNS infections, 10–15% to maternal-fetal infections and the remaining to various localized infections (compiled data from the French National surveillance system in the past 5 years). After a decrease in the number of cases in the second part of the 20th century, which correlates with the implementation of controls in food industry and information campaigns directed to pregnant women,^{11,12} the incidence of *Lm* has slightly re-increased in recent years, notably in Europe.^{8,13–16} The reason for this increase is unknown, but is likely a combination of the relative increase of the population at risk for listeriosis, such as the immunosuppressed host and/or elderly, and changes in food processing and habits.

The intracellular life of *Lm* and immune responses to this bacterium have been extensively studied in vitro and in vivo,^{17,18} yet many aspects of the pathophysiology of listeriosis, and particularly that of neuroinfection remain elusive, in part due to a relative lack of relevant and easy-to-handle animal model that reproduces all the hallmarks of the human disease (see below). Despite these limitations, epidemiological and clinical studies in human and ruminant as well as in vitro and in vivo mouse experimental infections, allow making a general picture of the putative mechanisms of *Lm* CNS invasion and infection. These studies suggest that diverse pathways could be used by *Lm* to gain access to the CNS and could include a retrograde neural route of invasion and crossing of the blood-brain barrier (BBB) by blood-borne bacteria.^{19,20} Yet, the molecular factors on both bacterial and host sides remain to be discovered, and new animal models recently developed will likely be instrumental to study and better understand the molecular mechanisms underlying neuroinfection (see below).

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Natural History of Neurolisteriosis

Lm and *L. ivanovii* are the two species of the genus *Listeria* (that includes *Lm*, *L. ivanovii*, *L. innocua*, *L. welschimeri*, *L. seeligeri*, *L. grayi*, *L. marthii* and *L. rocourtiae*) that have been described as pathogenic.⁷ *L. ivanovii* is almost exclusively recovered from ruminant, yet it can rarely infect humans²¹ and has not been described to disseminate to the CNS, while *Lm* is the only one that leads to CNS infection, in humans and in domestic ruminants,^{19,20} as well as in wild or domestic species such as cervidae (fallow deer²²), camelidae (llama²³), rodents (chinchilla and squirrels²⁴) and to a lesser extent carnivores: felidae (cat and cougar^{25,26}) and canidae (dog, fox and racoon dog^{24,27}). *Lm*-associated CNS infection can manifest as meningitis, meningoencephalitis, brain abscesses and rhombencephalitis,²⁸ the latter being typically associated with this bacterium. The ability of *Lm* to cause both acute meningitis and brain parenchymal infection differentiates it from other bacteria frequently responsible for meningitis such as *Streptococcus pneumoniae*,²⁹ *Neisseria meningitidis*³⁰ and *Haemophilus influenzae*,³¹ and underlines a commonality with *Mycobacterium tuberculosis*.³²

Of note, CNS infection in both humans and ruminants classically does not occur in pregnant individuals (data from the French National surveillance system and ref. 20), suggesting that (1) pregnancy is not a predisposing factor for neurolisteriosis and (2) placental and brain infection mechanisms are likely different. Interestingly, other human pathogens with a mucosal portal of entry, such as the apicomplex *Toxoplasma gondii*,³³ the bacterium *Treponema pallidum*³⁴ and enteroviruses,³⁵ have in common, as *Lm*, to disseminate to the CNS and the fetoplacental unit.

Neurolisteriosis in humans. CNS infection manifests in human as meningitis and meningoencephalitis, which are the most frequent clinical presentations observed in human, followed by brain stem infection (rhombencephalitis) and brain abscesses.

Meningitis and meningoencephalitis. Meningitis is defined as an inflammation of meninges, the protective membranes covering the central nervous system at the brain parenchyma and spinal cord levels. They are composed of the dura mater, the arachnoid and the pia mater. Meninges contain the cerebrospinal fluid, which is located in the subarachnoidal space, and vessels and capillaries that enter the brain parenchyma (Fig. 1).

Meningoencephalitis is a brain inflammation associated with meningitis. Infection of the meninges might either occur upon the crossing of the blood-meningeal barriers or as a consequence of bacterial dissemination to and from the brain parenchyma.

Meningitis and meningoencephalitis account for the majority of CNS infection by *Lm* in human (70 to 97%).^{36,37} *Lm* has been recognized as the second to fourth cause of community-acquired acute bacterial meningitis in adults.^{38,39} Known predisposing factors are immunosuppression, age over 50 y and underlying conditions such as malignancy or diabetes.^{36,37}

The clinical features of *Lm* meningitis differ from those of other bacterial meningitides in that *Lm* meningitis may have a sub-acute course (usually more than 24 h of symptoms before admission in hospital) and be associated with abnormal movements, seizures

and alteration of consciousness. The onset of such symptoms is evocative of dissemination to the brain parenchyma and their association with meningeal signs of meningoencephalitis. In a recent prospective study on bacterial meningitis in adults,³⁶ Brouwer et al. compared meningitis due to *Lm* and to other bacterial meningitis. Only patients with a positive CSF bacterial culture were included. This study showed that the classic triad of fever, neck stiffness and change in mental status was present in 13 (43%) of 30 *Lm*-infected patients, which was not significantly different to what was observed for other bacterial meningitis in the same cohort.³⁹ However, typical cerebrospinal fluid findings predictive for bacterial meningitis were frequently absent: up to 23% of patients had no CSF abnormality indicative of bacterial meningitis, compared with only 12% for other bacterial meningitis. Gram staining of cerebrospinal fluid samples revealed the causative organism in only 24% of patients, compared with 80% for other bacterial meningitis. This result is in agreement with other studies showing that Gram staining of the CSF is often negative,²⁸ reflecting a quantitatively modest dissemination of *Lm* in the CSF. Besides clinical descriptions, few data are available on *Lm*-associated meningoencephalitis. CT scan resolution has proven so far to be insufficiently precise to gain insight into the neuropathology of *Lm*-associated encephalitis, and magnetic resonance imaging (MRI) allows visualization of diffuse but so far rather unspecific lesions.^{40,41}

Brain abscess. In a small proportion of *Lm* CNS infections (up to 10%, with half of them in immunocompromised patients), macroscopic brain abscesses without meningeal involvement are observed.⁴² *Lm* brain abscesses are revealed by imaging (CT scan and/or MRI) and are preferentially located in subcortical areas, thalamus, pons or medulla. These unusual locations have been reported as evocative of their listerial origin.^{43,44} Less than 10% of these patients have no known underlying condition. This disease is also sub-acute, and CSF cultures are positive in 25% of cases, indicative of concomitant and possibly secondary meningitis.

Rhombencephalitis. Rhombencephalitis (i.e., brainstem encephalitis) is a pathology typically associated with *Lm*, and was first described in human by Eck in 1957.⁴⁵ Rhombencephalitis is characterized by progressive brainstem dysfunction. Clinical signs classically appear in two phases. In the first 4 to 10 d period, unspecific symptoms consisting in headache, malaise, nausea and vomiting can be present. In a subsequent period, asymmetrical cranial nerve deficits, cerebellar signs and hemiparesis or hemisensory defects, appear and can be concomitant to meningeal signs. Contrary to meningitis, rhombencephalitis occurs mostly in otherwise healthy people.^{46,47} Based on neuropathological analysis, it has been shown that these clinical signs correlate with inflammatory infiltrates predominantly present within nuclei, tracts and intraparenchymal parts of cranial nerves innervating the oropharynx (5th, 7th, 9th, 10th and 12th cranial nerves)⁴⁸ (see Fig. 2). These sequential observations in human may indicate that cranial nerve is secondarily reached via the extension of the infection process from the rhombencephalon, rather than a route for the retrograde transport of *Lm* to the rhombencephalon.

Neurolisteriosis in ruminants. Listeriosis is a common pathology in ruminants. Infection is supposed to occur mainly from

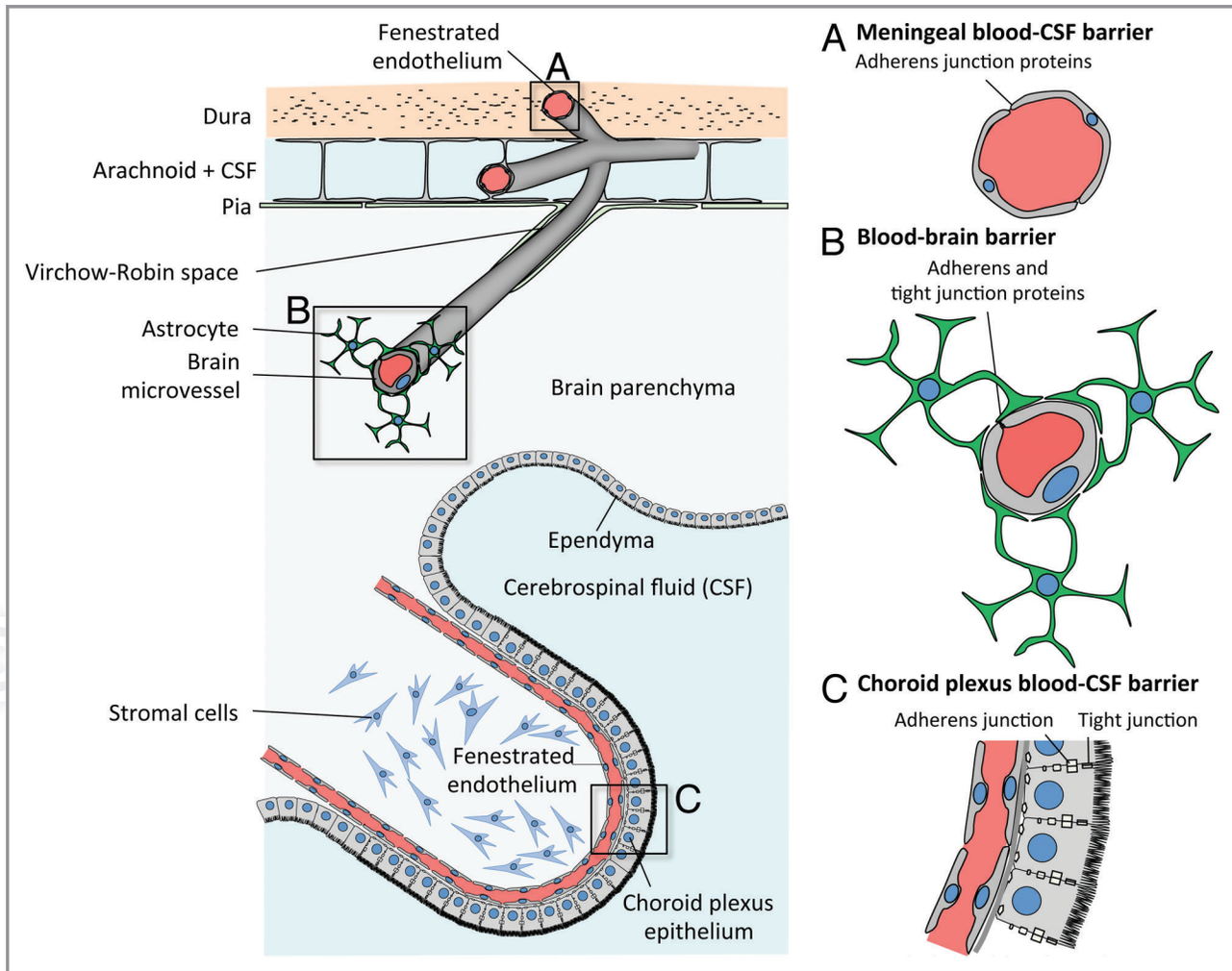


Figure 1. Blood-CSF and blood-brain barriers. From top to bottom: (A) Capillaries from the meningeal compartment, which represent part of the blood-CSF barrier. Dural endothelial cells are fenestrated, while endothelial cells from the subarachnoid space are joined by tight junctions. (B) Blood-brain barrier is made of endothelial cells joined by tight junctions. They are in close contact with astrocyte's "feet" (astrocyte's cell projection), which enable the differentiation of brain endothelial cell junctions. (C) Epithelial cells from the choroid plexus produce the cerebrospinal fluid (CSF) and isolate the fenestrated endothelium from the CSF, hence forming together with the meningeal endothelium the blood-CSF barrier. (Adapted from ref. 64.)

absorption of contaminated silage and manifests as abortion and encephalitis. Diffuse meningitis and meningoencephalitis are very rare in ruminants, contrary to rhombencephalitis, which is the cardinal form of disease, and was first described as "circling disease" in sheep in New Zealand.⁴⁹ A recent neuropathological survey in Switzerland identified *Neurospira* as the most prevalent CNS disease of small ruminants. The incidence of *Neurospira* in this country ranges from 200 to 500 cases/million small ruminants, very significantly exceeding the incidence in human. Of note, and as in human, placental infection and CNS infection seem to be two distinct pathologies, as concomitant infection of placenta and CNS is classically not observed in ruminant.²⁰

As in human, rhombencephalitis seems to occur mainly in hosts without known predisposing factor. The incubation period is long, from 1 to 7 weeks.^{50,51} Clinical signs, such as "circling disease," are linked to the CNS lesions, which are unilateral or

bilateral brainstem and cranial nerve deficits.^{52,53} The cranial nerves mostly involved are the 5th cranial nerve (trigeminal), 7th (facialis) and 12th (hypoglossus), and to a lesser extent the 6th, 8th, 9th and 10th nerves.⁵⁴ The prominence of cranial nerve deficit in ruminant, and the characteristics of their feeding behavior make of the retrograde transport of *Lm* to the rhombencephalon an attractive hypothesis, which is supported by neuropathological findings (see below).

Pathophysiology of *Neurospira*

What can be learned from *Lm* interactions with other host barriers? We have studied the molecular mechanisms underlying *Lm* ability to target and cross the intestinal and placental barriers in detail. We have shown that the *Lm* surface protein InA mediates a species-specific interaction with its host receptor E-cadherin (Ecad) at the intestinal epithelium level, resulting in

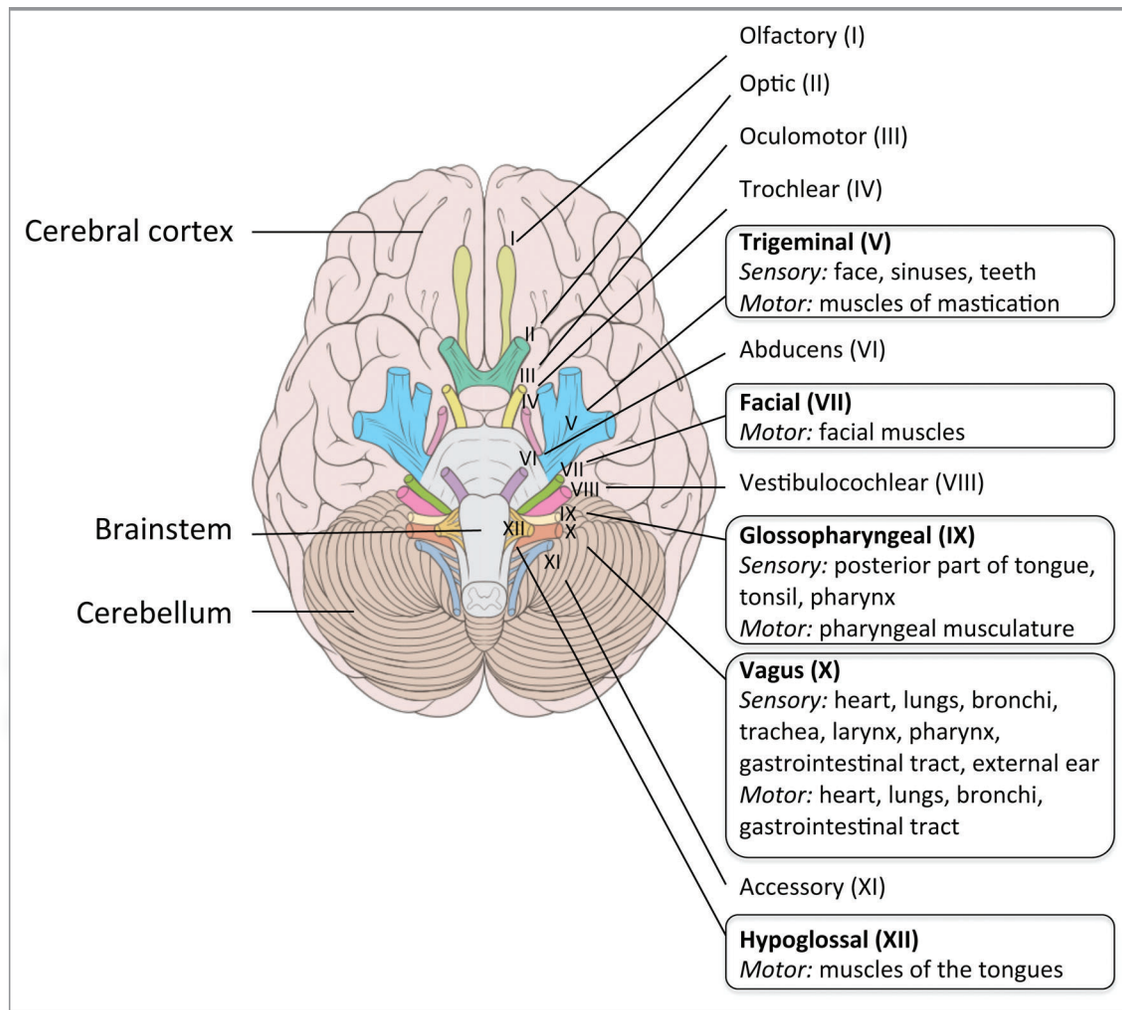


Figure 2. Cranial nerves in human. A frame surrounds the nerves that emerge from the regions most frequently infected by *Lm* in the brainstem. Adapted from Patrick Lynch; Creative Commons Attribution 2.5 License 2006; www.patricklynch.net.

bacterial internalization in enterocytes and crossing of the intestinal barrier.⁵⁵ Early after contamination, *Lm* crosses this barrier via transcytosis in mucus producing cells and their neighbors, which express lumenally accessible Ecad.⁵⁶ In contrast, InlB, another species-specific internalin of *Lm* that interacts with Met, is not involved in this process.^{57,58}

We have also shown that *Lm* targeting and crossing of the maternal-fetal (MF) barrier is critically dependent upon InlA interaction with trophoblast Ecad, but this time only in conjunction with InlB interaction with Met.^{57,59} The strong requirement of InlA in *Lm* crossing of the intestinal and placental barriers has been independently shown epidemiologically, since *Lm* isolates expressing a functional InlA are very significantly associated with their clinical origin, and particularly their fetoplacental origin.^{57,60}

These results demonstrate that the molecular mechanisms underlying *Lm* ability to target and cross the intestinal and MF barriers exhibit similarities but also critical differences. Remarkably, the BBB is also composed of Ecad- and Met-expressing cells, such as the microvascular endothelium and choroid plexus

epithelium. Thus, the molecular mechanisms enabling *Lm* to cross the BBB may also implicate InlA and/or InlB, and thus also share similarities with *Lm* crossing of the intestinal and MF barriers.

Crossing of the blood-brain, the blood-CSF barriers, or both? *Lm*-associated meningoencephalitis and rhombencephalitis appear to be two distinct forms of neurolisterosis. Yet, it is not established whether the mechanisms underlying these two pathologies actually differ or share common steps. The hematogenous route of infection is favored for meningoencephalitis, whereas the neural retrograde route, described thereafter, has been proposed for *Lm*-associated rhombencephalitis in ruminants. Two major and non-exclusive ways for bacteria to cross the BBB are proposed, and described in **Figure 3**: extracellular bacteria, either free in the blood and/or associated to cells, may recognize receptors at the surface of the barriers and cross them (**Fig. 3A**); alternatively, and since *Lm* is a facultative intracellular bacterium, this bacterium may gain the CNS in infected cells, such as circulating leucocytes, which are known to be able to cross themselves the BBB (**Fig. 3B**).⁶¹

Brain endothelial cells and choroid plexus epithelial cells form the blood-CNS barriers. Brain endothelial cells that constitute the blood-brain barrier differ in many ways from classical endothelial cells.⁶² They are in contact with each other by tight junctions, made of occludin, claudins, junctional adhesion molecules (JAM) and cell-selective adhesion molecules that induce a high trans-endothelial resistance and prevent paracellular trafficking of macromolecules. In addition they express adherens junction proteins such as VE-cadherin and N-cadherin (Fig. 1). Rubin et al. have also proposed that brain endothelial cells express E-cadherin, although this result would need to be revisited with updated tools, as they predate the discovery of VE-cadherin.⁶³ Differentiation of these cells is allowed and maintained by an intimate interaction with astrocytes and their secreted products and neurons in what is referred as “neurovascular unit.”⁶⁴

Choroid plexus epithelial cells form the blood-CSF barrier. They are located in brain ventricles and produce CSF from a blood filtrate.⁶⁵ They are separated from the systemic circulation by fenestrated endothelial cells filled with blood and stromal cells (Fig. 1). As other epithelial cells, they express E-cadherin at their

basolateral pole, as well as tight junctions and as such constitute an attractive site of entry for *Lm* in the CNS.

Direct interaction of extracellular bacteria with the blood-brain and blood-CSF barriers. As mentioned before, *Lm* has the capacity to interact via InlA and InlB with its cellular receptors E-cadherin and Met, respectively, to cross the intestinal and placental barriers. Since these two receptors are expressed at the surface of choroid plexus epithelial cells, and Met is also expressed at the brain endothelial level, similar mechanisms than at the intestinal and placental levels could occur at the blood-CSF and BBB, respectively.

It has been shown in vitro that *Lm* is able to invade human umbilical vein endothelial cells (HUVEC) in an InlB-dependent manner,⁶⁶ as well as human brain microvascular endothelial cells (HBMEC).⁶⁷ In these cells, bacteria can survive, replicate, move intracellularly by inducing actin tail formation and spread from cell to cell.^{67,68} However, widespread vasculitis is not observed in *Lm* infection and interaction with HBMEC may result in vivo in a rapid crossing of the barrier rather than local infection and replication in brain microvessel endothelium.

Another bacterial candidate is *Vip*, which can interact with its receptor the heat-shock protein Gp96 expressed at the brain microvessel surface.⁶⁹ It has been shown that Δvip strain is attenuated in the brain invasion upon intravenous inoculation, but also in other organs such as liver and spleen, which may indeed account for the attenuated CNS infection observed with the Δvip deletion mutant. Of note, Gp96 has been shown to be a potential receptor for another pathogen targeting the brain, *Candida albicans*,⁷⁰ while a related protein, Ec-Gp96, may also be involved in the targeting of HBMEC by *E. coli* K1, a prominent cause of neonatal meningitis.⁷¹

Lm is able to induce, via the action of the pore forming toxin listeriolysin O (LLO) NF κ B activation, both in cultured endothelial cells and in brain microvessels, and induce the expression of the surface adhesion molecules P- and E-selectin, ICAM-1 and V-CAM-1, as well as interleukin (IL) IL-6 and IL-8 and the chemoattractant MCP-1, which allow neutrophil and monocyte adhesion to the endothelial cells.⁷²⁻⁷⁴ These effects may modulate blood-brain barrier function and favor *Lm* access to the CNS.

Trojan horse model. The “Trojan horse” mechanism was first proposed for ovine CNS invasion by visna virus.⁷⁵ *Lm* is able to escape from the phagosome in a LLO-dependent manner and to move in the host cytosol in an ActA-dependent manner.¹⁸ These properties enable *Lm* to survive and proliferate in phagocytic cells such as monocytes or dendritic cells both in vitro and in vivo.^{76,77} Infected phagocytic cells may translocate, as a Trojan horse, *Lm* into the CNS. In mice intravenously inoculated, *Lm* has been shown to infect the brain even after gentamicin treatment, which is thought to eradicate extracellular blood-borne bacteria.¹⁹ This suggests that infected phagocytes may play a role in brain invasion. Further experiments showed that the main infected cells were Ly-6C^{high} monocytes, which could be isolated from infected brains.⁷⁸ In another study of acute infection in mouse, it has been observed that bacteria invade bone marrow myelomonocytic cells expressing the phenotype

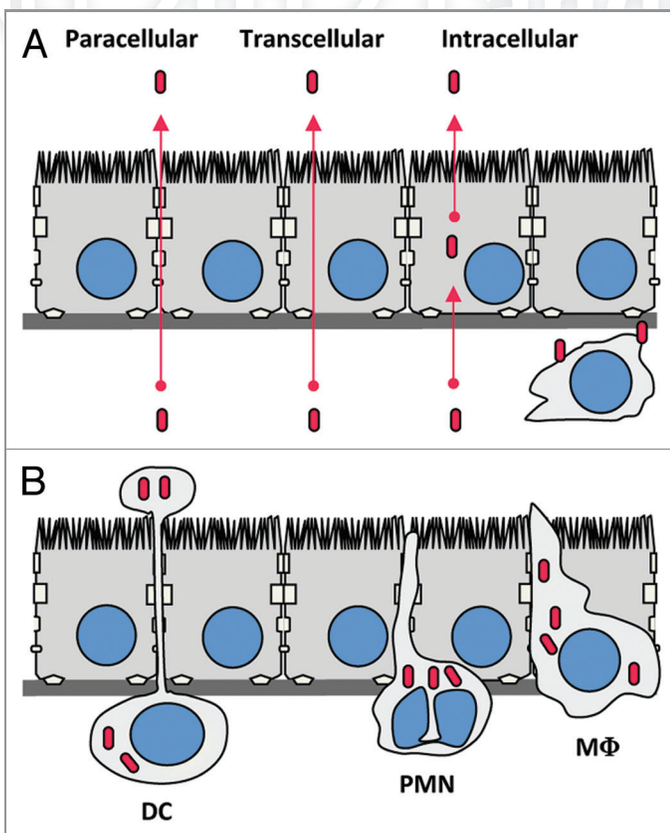


Figure 3. Potential mechanisms by which *Lm* could cross the blood-brain barrier. (A) Extracellular *Lm*, either free in the blood and/or associated to circulating cells, may recognize receptors at the surface of the barriers (as InlA, InlB or Vip) and cross them. (B) Trojan horse mechanism: Circulating leucocytes infected by *Lm*, such as monocytes, dendritic cells or polymorphonuclear cells, may cross the BBB hence targeting the bacteria in the CNS. DC, dendritic cell; PMN, polymorphonuclear leukocyte; MΦ, monocyte/macrophage.

CD31^{pos};Ly-6C^{pos};CD11b^{pos};LY-6G^{low}.⁷⁹ Injection of *Lm*-infected bone marrow cells appeared to facilitate CNS invasion in comparison to free bacteria or infected splenocytes, indicating that these infected cells may disseminate to the CNS. Yet the observation of a concomitant higher bacterial load in blood probably rather indicates that they constitute a niche in which bacteria can survive and multiply. The mechanisms by which these cells target the CNS is not known, although this process has been shown as IFN γ -dependent but CCR2-independent.⁸⁰ In the infected host, the action of LLO on endothelial cells may also lead to ICAM-1 and P-selectin expression, which in turn would allow infected cells to adhere to brain microvessels or meningeal vessels and favor CNS invasion.^{72,81}

When circulating in the brain vasculature, infected cells could translocate from the blood to the CNS. Direct neuronal infection is not a necessary step to induce brain damage upon infection. It has been observed that neurons are not easily directly infected, yet in vitro observations indicate that they could be infected indirectly by bacteria spreading from infected macrophages to neurons.⁸² Bacteria may also spread from myeloid infected cells to endothelial cells and thus cross the BBB, although this has only been observed in vitro.⁶⁷ This phenotype has been shown to depend on both ActA (which mediates actin-based motility that result in intracytosolic movement and formation of bacteria containing protrusions able to penetrate neighboring cells) and on phospholipase C PlcA and PlcB, (which are implicated in the lysis of the double membrane vacuole in the recipient cell).⁶⁷

Neural retrograde route. In ruminants and more generally in the process of rhombencephalitis, it has been proposed that infection via neural retrograde transport is the most likely to occur.²⁰ This hypothesis is based on the observation of the neuropathological pattern of natural disease. Oevermann et al. have reported the results of their detailed investigations on the neuropathology of more than 200 cases of *Lm*-associated encephalitis in cattle, sheep and goats from Switzerland.⁵⁴ They analyzed the anatomical distribution, severity and bacterial load and tried to define a posteriori the temporal evolution of the lesions. Their results suggest that *Lm* gains access to the brainstem via retrograde axonal migration along the trigeminal nerve branches and also along other cranial nerves. Moreover, *Lm* seems to spread further from the brainstem into rostral brain regions, likely by intracerebral axonal migration, as evidenced by histopathological analyses.

In an attempt to investigate in a murine model if *Lm* could induce rhombencephalitis via a retrograde neural route, bacteria were inoculated unilaterally into facial muscle, or peripheral parts of a cranial nerve. In this model, clinical and histological signs of mainly ipsilateral rhombencephalitis could be observed.⁸³ This suggests that rhombencephalitis could be caused by intra-axonal bacterial spread from peripheral sites to the central nervous system.

The way peripheral nerves become infected in the first place, prior to retrograde dissemination to the CNS remains unknown. It has been proposed that bacteria might enter through mucosal injury in the oropharyngeal cavity.²⁰ Interestingly, in human, case reports of rhombencephalitis have been described in patients after

dental surgery,^{84,85} supporting this hypothesis. Clinical signs of rhombencephalitis have also been observed in orally inoculated mice after a scarification of the oral, nasal and labial mucosae.⁵⁰

Recently it has been proposed that E-cadherin could act as a receptor for *Lm* to gain access to the brainstem.⁸⁶ InLA receptor Ecad is indeed expressed in satellite cells and myelinating Schwann cells in cranial nerves and ganglia, and *Lm* can be observed in phagocytes, axons, Schwann cells, satellite cells and ganglionic neurons in ruminant biopsies. The authors suggest that the oral epithelium and Schwann cells expressing E-cadherin may provide a portal of entry for free bacteria, which may then invade the axonal compartment by cell-to-cell spread. To prove this hypothesis, an experimental model of rhombencephalitis in which the role of InLA could be tested is needed. We are currently working on this.

Schluter et al. demonstrated that once in the brain after intracerebral infection, InLA and InLB play no role in a mouse model, but PlcB, which is involved in vacuolar escape and cell-to-cell spread does.⁸⁷ Although we now know that InLA is inefficient in the mouse,⁸⁸ these results are in favor of an infection of neurons by a spread from cell to cell.

Which Bacterial Factor(s) May Be Implicated in *Lm* CNS Tropism?

Epidemiological analyses. We have shown that clinical samples express a full-length InLA far more frequently (96%) than strains recovered from food samples in which 35% of strains harbors a truncated InLA. This accounts for the role of InLA in crossing the intestinal barrier.⁶⁰ Full-length InLA is also statistically associated with maternofetal listeriosis compared with bacteremia, this again showing the role of InLA in invading the placenta and fetus.^{57,60} However, even if amount of full-length InLA strains in neuro-listeriosis sample (98%) is higher than those from bacteremia (93%), this difference does not reach statistical significance. So far, involvement of InLA in inducing neuro-listeriosis is unclear, even if it could play a role in rhombencephalitis (see above). Experiments in permissive animal models for InLA-E-cadherin interaction are under way and will allow knowing precisely the role of InLA in *Lm* CNS tropism in vivo.

In an attempt to find factors involved in neuro-listeriosis, a multi-locus variable-number tandem-repeat analysis (MLVA) has been conducted on *Lm* isolates from ruminant rhombencephalitis and were compared with isolates from human patients, food and environment.⁸⁹ Allelic profile-based comparisons grouped *Lm* mainly into three clonal complexes. Isolates from human and ruminant brain samples were mainly located in clonal complex A, which contains all but one rhombencephalitis isolate from cattle. A larger study is now needed to compare isolates from CNS invasion to isolates from bacteremia in order to determine genes specifically involved in CNS invasion.

High-throughput methods. In order to determine genes that could be involved in CNS invasion in a mouse model of infection, Autret et al. induced brain infection combined with signature-tagged mutagenesis (STM),⁹⁰ and identified genes encoding cell wall components such as *gtcA*, which is involved

in the glycosylation of teichoic acid, and *ytgP*, coding for a putative integral membrane protein possibly involved in polysaccharide biosynthesis. However, in a second STM experiment aimed at discovering genes involved in the survival in liver, the same *gtcA* and *ytgP* mutants were found to be attenuated,⁹¹ suggesting a non-specialized role for these genes in the CNS invasion, which could be involved in the overall in vivo proliferation or survival of the bacterium.

New Animal Models are Needed to Study Neurolisteriosis

Even if advances in the pathophysiology of neurolisteriosis have been achieved from epidemiological analyses in human and ruminants, as well as from observations of naturally infected ruminant or experimentally infected mice, many questions remain about the different routes of CNS invasion. There is now a need for a small animal model of neurolisteriosis that would recapitulate the different steps and features of the CNS infection in human.

This model has to possess the following properties (1) to be, as in human, permissive for the known virulence factors InIA and InIB, (2) to develop the clinical hallmarks of human neurolisteriosis (meningitis and/or rhombencephalitis) and (3) to exhibit the neuroradiological and histopathological lesions observed in humans.

Mouse has been proven as an animal model of choice for studying immunity and the role of virulence factors of *Lm* after an intravenous injection, such as listeriolysin or actA. Different approaches have been performed to induce neurolisteriosis in mice. For all of them, one of the concerns is that mouse E-cadherin is not recognized by InIA, contrary to human E-cadherin, thereby questioning the relevance of results obtained in this model.^{88,92} A classical approach is to inject *Lm* in the tail vein. This method is adapted to study meningitis if bacteria cross the BBB from the bloodstream in an InIA-independent manner.⁷⁹ However, since *Lm* is a foodborne pathogen, this route of infection bypasses a critical stage of the infection, the crossing of the oro-intestinal barrier, which may affect the way by which bacteria access the bloodstream.

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In order to study CNS infection in small animal models, rodents permissive to both InIA and InIB, such as gerbil and humanized mice could be used.⁹³ A first assay has been conducted in gerbils by Blanot et al.⁹⁴ However, bacteria where injected in the middle ear, a model highly relevant for CNS infections that stems from a contiguous infection of the upper airway apparatus (such as pneumococcal and hemophilus-associated meningitis) but which relevance is uncertain for a foodborne infection.

Of note, gerbil and humanized mice are permissive to InIA and InIB but, like any animal model, it cannot be excluded that other host-bacterial interactions that occur in human do not occur in one of these rodents. The reverse is true and interactions that occur in a given animal species may not occur in human. It is therefore important to study listeriosis in various animal species and to compare clinical, microbiological, neuro-imaging data with what observed in human, in order to fully understand the pathophysiology of neurolisteriosis, and uncover its potential species-specific characteristics.

Conclusions

Neurolisteriosis is a major health and economical concern. Deciphering the mechanisms by which *Lm* gains access to the brain will help us not only understand a critical virulence attribute of *Lm* but also hopefully help improve the prognosis of this still very severe infection. *Lm* is a model to understand how pathogens cross host barriers, and studying neurolisteriosis and identifying novel bacterial and host factors implicated in the development of neurolisteriosis will likely be beneficial for a better understanding of the basic mechanisms underlying CNS infections.

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