Ovine Proinflammatory Cytokines Cross the Murine Blood-Brain Barrier by a Common Saturable Transport Mechanism

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Abstract
Objectives: The cytokines interleukin (IL)-1\textbeta and IL-6 are modulators of the neuroimmune axis and have been implicated in neuronal cell death cascades after ischemia or infection. Previous work has shown that some cross-species conservation exists between human and rodent blood-brain barrier (BBB) transport systems. To further assess cross-species conservation of cytokine transport across the BBB, the current studies investigated permeability and inhibition of ovine IL-1\textbeta and IL-6 in the mouse. Methods: IL-1\textbeta or IL-6 was radioactively labeled with \textsuperscript{131}I and injected into the jugular vein at time zero. A subset of mice received 1 or 3\textmu g/mouse of an unlabeled ovine or murine cytokine (IL-1\textbeta or IL-6) to assess self- and/or cross-inhibition of transport. Permeability was assessed using multiple-regression analysis. Results: There was a significant linear relationship for both ovine \textsuperscript{131}I-IL-1\textbeta and \textsuperscript{131}I-IL-6 between brain/serum ratios and exposure time, indicating BBB permeability. Inclusion of 3\textmu g/mouse unlabeled ovine IL-1\textbeta or IL-6 significantly reduced the transport of ovine \textsuperscript{131}I-IL-1\textbeta or \textsuperscript{131}I-IL-6, respectively, across the BBB. Transport of both ovine \textsuperscript{131}I-IL-1\textbeta and \textsuperscript{131}I-IL-6 was significantly inhibited by 1\textmu g/mouse of murine IL-1\textbeta or IL-6, respectively. In contrast, 1\textmu g/mouse of unlabeled ovine IL-1\textbeta or IL-6 did not inhibit the transport of murine \textsuperscript{131}I-IL-1\textbeta or \textsuperscript{131}I-IL-6. Conclusions: Ovine IL-1\textbeta and IL-6 cross the mouse BBB by saturable transport. Inhibition of transport by murine homologs indicates that both species use the same transport mechanisms. Conversely, an inability of ovine cytokines to significantly inhibit the transport of murine cytokines indicates that mouse BBB has a lower affinity for ovine than murine cytokines. Knowledge of species-conserved BBB transport mechanisms may facilitate the development of novel animal models of central nervous system pathogenesis.

Introduction
Neuroimmune interactions have gained increasing attention as research focused on neurodegenerative pathogenesis, and brain injury continues to implicate immune responses and cytokine upregulation in a variety of central nervous system (CNS) disorders including stroke,
multiple sclerosis and infection [1–7]. Despite historical doubts about the transport and interaction of peripherally derived cytokines with CNS function and disease, research continues to reveal mechanisms for cytokine transport across the intact and damaged blood-brain barrier (BBB) [7–10]. Proinflammatory cytokines such as interleukin-1β (IL-1β) and interleukin-6 (IL-6) have been shown to cross the murine BBB by saturable transport systems [8–11]. This saturable transport involves nonionic-driven protein-mediated transfer across the BBB [1]. Importantly, despite the fact that the BBB impedes leakage of molecules into the brain, it also serves as an interface for regulatory proteins [1]. Cytokines are one such family of regulatory proteins, with a diverse set of functions, which have been implicated in normal and pathological conditions [1, 12]. A number of animal models continue to be used to address critical questions regarding species-conserved mechanisms of cytokine transport through the BBB, with potential implications for human neurological disorders and pathological modeling.

Saturable BBB transport systems have been identified for a number of murine cytokines including, IL-1β, IL-1α, IL-6 and tumor necrosis factor-α (TNF-α). Further, cross-species studies have shown some conservation of these mechanisms. For example, human IL-6 and IL-1α cross the murine BBB by saturable transport although with less affinity than the murine homologs, which share 60–70% amino acid identity to human ILs [13–15]. However, these mechanisms were not conserved for human IL-1β and TNF-α when assessed for murine BBB permeability [16, 17]. Although a number of different species, including murine and ovine, have been used independently to examine BBB transport mechanisms and general permeability across ontogeny [1, 18, 19], cross-species studies continue to be undertaken to assess phylogenetically conserved transport systems. The mouse represents a highly efficacious model for assessing BBB permeability from a logistical standpoint given the relatively low cost to achieve statistical power. Nonetheless, kinetic studies of BBB function cannot be performed in murine fetuses or newborns because of size limitations and because vascular access is not feasible.

Recent findings suggest that proinflammatory cytokines are important in the pathogenesis of brain damage in premature human infants [20–25]. Further, evidence also suggests that white matter lesions produced in the ovine fetus are similar to the lesions of periventricular leukomalacia, which represent an important antecedent of cerebral palsy in premature infants [26–28]. These findings [26, 29] collectively suggest that the preterm ovine fetus is an excellent model for the brain damage observed in premature human infants [26, 29–31]. Therefore, models of neuroinflammation related to cytokine upregulation and BBB permeability in the ovine fetus may represent a more ethological model of human in utero pathogenesis because of the ovine’s extended gestational length, ability for real-time physiological monitoring and the frequency of single fetuses as compared with the mouse [18, 19, 26, 29, 32]. Therefore, complementary cross-species studies assessing BBB transport mechanisms in mice – before more detailed studies in ovine models – may shed light on evolutionarily conserved systems and provide a steppingstone for the development of ethologically relevant models of neuropathogenesis related to the regulation of proinflammatory cytokines.

Although murine and ovine IL-1β and IL-6 share approximately 58% and 40% identity, respectively, based on basic local alignment search tools [33, 34], the ability of human ILs to cross the murine BBB and bind to the same receptors suggests species-conserved mechanisms despite variation in protein homology [14, 15, 35]. Therefore, we assessed the ability of radioactively iodinated ovine IL-1β (131I-IL-1β) and (131I-IL-6) to enter the murine brain after intravenous injection. We further assessed the ability of ovine and murine IL-1β and IL-6 to self- and cross-inhibit the entry of 131I-IL-6 and 131I-IL-1β into the murine brain. These studies may facilitate the development of novel animal models of CNS pathogenesis related to BBB permeability.

**Materials and Methods**

**Subjects**

Subjects were 8- to 12-week-old male CD-1 mice from our in-house colony. Mice were provided food and water ad libitum. All procedures and protocols comply with the National Institutes of Health guide for care and use of laboratory animals. All procedures were approved by the Institutional Animal Care and Use Committee of Veterans Affairs Medical Center of Saint Louis and the Saint Louis University School of Medicine.

**Purification and Iodination of Cytokines**

Ovine IL-1β and IL-6 were obtained from Commonwealth Scientific and Industrial Research Organization (CSIRO) Animal Health (Parkville, Vic., Australia) that had been cultivated in certified (virus-free) culture broths. The bacterial samples were lysed, sonicated, the proteins column-purified, and then sterile filtered to remove any possibility of bacterial contamination as previously described (for IL-1β [36, 37] and IL-6 [38]). The purity of ovine IL-6 was 95% and that of IL-1β was 90% as evidenced by SDS-PAGE and Western immunoblot analysis, consistent with that described by McWaters et al. [38] and Rothel et al. [36], re-
spectively, using the same methods. Murine cytokines (IL-1β and IL-6) were obtained from R&D Systems (Minneapolis, Minn., USA). The IL-1βs were labeled with 131I by the chloramine-T method and purified on a column of Sephadex G-10 as previously applied to IL-1 [14]. Briefly, the iodinated IL-1βs were separated from free iodine on a column of G-10 Sephadex that had been hydrated in 0.25 M phosphate buffer with 1% BSA. Prior to separation of the iodination mixture, the column was washed free of excess BSA with phosphate buffer, which was also used to elute the radiolabeled ILs. The IL-6s were labeled with 131I by the lactoperoxidase method as previously described [39]. Briefly, IL-6s were mixed with a solution of 0.4 M sodium acetate (pH 5.6), lactoperoxidase (10 µg/ml), and 1 mCi of 131I and purified with high performance liquid chromatography. Chloramine-T was used for IL-1β radiolabeling because of its higher labeling efficiency as compared to the lactoperoxidase method. However, IL-6 was labeled with lactoperoxidase given that IL-6 does not remain biologically active following chloramine-T-mediated radiolabeling.

Measurement of Influx Rate

Subjects were anesthetized with urethane and the right jugular vein and left carotid artery were exposed as previously described [39]. Five micrograms of cytokine (IL-1β or IL-6) was radioactively labeled with 1 mCi of 131I. Label integrity was evaluated using acid precipitation. If activity decreased to less than 90% of the original level, the material was excluded from the study. A volume of 0.2 ml of Ringer’s solution containing 1% by weight of BSA and 3 × 10^6 counts per minute (cpm) were injected into the jugular vein at time zero as previously described [39]. Blood was collected from the right carotid artery and the animals were decapitated between 1–15 min after the injection. Subsequently, the levels of radioactivity in brain and blood samples were measured in a gamma counter. A subset of mice received 1 or 3 µg/mouse of an unlabeled ovine or murine cytokine (IL-1β or IL-6) included simultaneously with the labeled protein in order to assess affinity and/or cross-inhibition of transport. Cross-inhibition studies similar to those previously described [14] were conducted to determine whether the transporter for ovine IL-1β was the same as that previously identified for murine IL-1β. The unidirectional influx rate from blood to brain (K_i, in µl/g·min) and the initial volume of distribution in brain at time zero (V_i, in µl/g) were determined by multiple-time regression analysis. Importantly, V_i accounts for the amount of radioactivity in the vascular space and endothelial receptor binding at time zero allowing for accurate calculation and correction of K_i. The brain/serum ratios (in µl/g) were plotted against exposure time (Expt, in min):

\[ \text{Am/Cpt} = K_i(\text{Expt}) + V_i \]

where Am is the cpm/g of brain, Cpt is the cpm/µl of serum at time (t), and

\[ \text{Expt} = \int_0^t \text{Cpt}(t)dt/Cpt \]

where t is the time and dt is a dummy variable for time. Use of exposure time corrects for clearance of 131I-ILs from blood, mathematically creating a steady-state for the serum levels. The linear portion of the relation between brain/serum ratios and Expt is used to calculate K_i. Regression lines were compared statistically with the GraphPad Prism 5.0 program (GraphPad Inc., San Diego, Calif., USA).

Results

IL-1β Permeability

Results revealed a significant linear relationship between brain-serum ratios and exposure time for ovine 131I-IL-1β. Each point represents 1 subject. Histogram shows that inclusion of 3 µg/mouse but not 1 µg/mouse of unlabeled ovine IL-1β significantly reduced the transport of ovine 131I-IL-1β across the BBB. Error bars are SEM. * p < 0.05.

IL-6 Permeability

Results revealed a significant linear relationship between brain-serum ratios and exposure time (Expt) (r =
0.96, n = 9, p < 0.001) for ovine $^{131}$I-IL-6 with the $K_i$ (slope) = 1.049 ± 0.123 µl/min (fig. 3a). Further, inclusion of 3 µg/mouse unlabeled ovine IL-6 (n = 10 with unlabeled, n = 10 without) significantly [F (1, 18) = 15.4, p < 0.001] reduced the transport of ovine $^{131}$I-IL-6 across the BBB, indicating a saturable transport mechanism (fig. 3b). Cross-inhibition studies showed that the transport of murine $^{131}$I-IL-6 was not inhibited by 1 µg/mouse of unlabeled ovine IL-1β (fig. 2a). Further, inclusion of 3 µg/mouse unlabeled ovine IL-6 (n = 10 with unlabeled, n = 10 without) significantly [F (1, 18) = 15.4, p < 0.001] reduced the transport of ovine $^{131}$I-IL-6 across the BBB, indicating a saturable transport mechanism (fig. 3b). Cross-inhibition studies showed that the transport of murine $^{131}$I-IL-6 was not inhibited by 1 µg/mouse of unlabeled ovine IL-6 (p = NS). In contrast, administration of 1 µg/mouse unlabeled murine IL-6 (n = 7 with unlabeled, n = 8 without) significantly [F (1, 13) = 4.67, p < 0.05] inhibited the transport of ovine $^{131}$I-IL-6 across the BBB.

**Discussion**

These studies show that both ovine IL-1β and IL-6 cross the BBB of the mouse by saturable transport. Specifically, results revealed a significant linear relationship between brain/serum ratios and exposure time for ovine $^{131}$I-IL-6 (fig. 3a). Further, inhibition of $^{131}$I-IL-1β (IL-1β and IL-6) transport by the murine homologs suggests competitive transport, and that ovine cytokines use the same transporters as those of the mouse. Importantly, inclusion of 3 µg/mouse, but not 1 µg/mouse, ovine IL-1β significantly reduced the transport of ovine $^{131}$I-IL-1β across the BBB. Moreover, inclusion of 3 µg/mouse ovine IL-6, but not 1 µg/mouse, significantly inhibited...
ovine 131I-IL-6 transport into the murine brain. In contrast, transport of ovine 131I-IL-1β and 131I-IL-6 was significantly inhibited by only 1 μg/mouse of murine IL-1β and IL-6, respectively. The inability of ovine cytokines to inhibit the transport of murine cytokines significantly indicates that the mouse BBB has a lower affinity for ovine than murine cytokines.

The results presented above identify common saturable BBB transport mechanisms for ovine and murine IL-1β and IL-6. Although the mode of transport appears to be conserved between species, ovine cytokines exhibit less affinity for transport as compared to their murine counterparts. These differences are likely to originate from clear disparities in amino acid homology, which may be reflected in the binding properties of the transport systems. Further, mouse ILs were able to inhibit entry of the ovine ILs, suggesting either multiple carriers with overlapping affinity or a single carrier able to transport both murine and ovine ILs across the BBB [14]. Regardless of the mechanism, it is clear that the murine system preferentially transports the native ILs over those of the ovine species. It should be emphasized that most cytokine transport systems are specific for closely related cytokines. For example, the IL-1 system is unable to transport TNF-α or IL-6 [9]. Further, across species, some cytokines have increased receptor selectivity and higher BBB uptake rates than others. For example, the mouse has a higher blood to brain transport rate of human 125I-IL-1α as compared to that of the rat [1, 17]. Although variations in transport rates and affinities can be seen across species, saturable transport for cytokines IL-1β and IL-6 appears to be conserved for both the ovine and murine proteins. Given that similar transport systems have been observed for human, murine and now ovine ILs [13, 14], the current study opens the door to novel translational models of proinflammatory cytokine-mediated pathogenesis in the ovine [19, 26, 29].

**Conclusions**

The current studies provide additional evidence that transport mechanisms for proinflammatory cytokines IL-1β and IL-6 are conserved across several mammalian species, including ovine. Further, the ability of ovine cytokines to enter the murine brain from blood by saturable transport provides further support for a strong link between the peripheral immune system and the CNS. Finally, the present findings provide a starting point for more in-depth investigations of BBB permeability and cytokine-mediated pathogenesis in highly valuable ovine models of human neurodevelopmental pathology.

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**References**


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