Influence of Enriched Environment on Viral Encephalitis Outcomes: Behavioral and Neuropathological Changes in Albino Swiss Mice

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**Abstract**

An enriched environment has previously been described as enhancing natural killer cell activity of recognizing and killing virally infected cells. However, the effects of environmental enrichment on behavioral changes in relation to virus clearance and the neuropathology of encephalitis have not been studied in detail. We tested the hypothesis that environmental enrichment leads to less CNS neuroinvasion and/or more rapid viral clearance in association with T cells without neuronal damage. Stereology-based estimates of activated microglia perineuronal nets and neurons in CA3 were correlated with behavioral changes in the Piry rhabdovirus model of encephalitis in the albino Swiss mouse. Two-month-old female mice maintained in impoverished (IE) or enriched environments (EE) for 3 months were behaviorally tested. After the tests, an equal volume of Piry virus (IEPy, EEPy)-infected or normal brain homogenates were nasally instilled. Eight days post-infection (dpi), when behavioral changes became apparent, brains were fixed and processed to detect viral antigens, activated microglia, perineuronal nets, and T lymphocytes by immuno- or histochemical reactions. At 20 or 40 dpi, the remaining animals were behaviorally tested and processed for the same markers. In IEPy mice, burrowing activity decreased and recovered earlier (8–10 dpi) than open field (20–40 dpi) but remained unaltered in the EEpy group. EEpy mice presented higher T-cell infiltration, less CNS cell infection by the virus and/or faster virus clearance, less microgliosis, and less damage to the extracellular matrix than IEPy. In both EEpy and IEPy animals, CA3 neuronal number remained unaltered. The results suggest that an enriched environment promotes a more effective immune response to clear CNS virus and not at the cost of CNS damage.

**Introduction**

Sublethal encephalitis following viral infections is known to affect behavior and the immune response, and viral diseases of the central nervous system (CNS) represent a significant proportion of neurological disabilities, particularly in poor countries [1]. Emerging virus infections of the CNS are mainly associated with RNA viruses and many that cause neurologic disease [2]. The rhabdoviruses are a part of the broad group of negative-strand RNA viruses, a group that includes a number of medically relevant viruses such as avian influenza, measles, Ebola, and vesicular stomatitis virus (VSV) [3]. Because VSV has limited human pathogenicity, it has been used as a model of rhabdoviruses in both in vitro and in vivo studies investigating viral adaptive and host immune responses [4,5].

An enriched environment (EE) was previously described as enhancing natural killer (NK) cell activity, including their recognition and killing of virally infected cells [6]. An enriched environment has been defined as social interactions with con-specifics and a stimulation of exploratory and motor behavior with a variety of toys, ladders, tunnels, rope, bridges, and running wheels for voluntary physical exercise changed periodically, as opposed to an impoverished environment with reduced to social interactions [7]. Rats reared under EE conditions present immune cell recruitment with a higher number of activated microglia than control rats, and these ramified microglial cells resemble the neuroprotective phenotype of microglia activated by T-cell–derived cytokines [8].

A variety of viruses on the surface of host cells target specialized features of the extracellular matrix, known as perineuronal nets

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(PNs) [9], which consist of glycosaminoglycans. Virus affinity for glycosaminoglycans is a determinant of cell tropism, loss of invasiveness, and reduced efficiency in viral spreading through the circulation [10,11,12]. Because environmental enrichment increases the number of PNs [13,14], of interest is whether or not viral spreading and neuroinvasion are reduced in animals housed under the EE condition.

Infections of the CNS with cytopathic neurotropic viruses, such as vesiculoviruses, require the parenchymal penetration of dendritic cells, T lymphocytes, and microglial activation for virus clearance and survival [15,16,17]. Because T cell migration to the brain parenchyma enhances viral clearance in VSV encephalitis [18], we tested the hypothesis that an EE may increase T cell migration to the parenchyma and promote faster virus clearance from the brain.

Thus, in the present report, we induced Piry viral encephalitis in an adult albino Swiss mouse model housed in an impoverished environment (IE) or EE to investigate the hypothesis that an EE may reduce neuropathological damage and behavioral changes and promote less CNS invasion and/or faster virus clearance from the brain. We found that compared to infected IE animals, infected EE animals presented less viral neuroinvasion, less microglial activation, less damage in the specialized extracellular matrix, greater infiltration of CD3-immunolabeled T-lymphocytes in the brain parenchyma, and reduced behavioral changes.

Results

Piry virus encephalitis, cell targets, and areas of neuroinvasion

Figure 1 illustrates the cellular targets and areas of neuroinvasion in adult female mice at 8 days post-instillation (dpi). Piry viral antigens in the brain parenchyma were revealed in the cytoplasm of infected cells that stained positive for virus proteins. This feature is consistent with the fact that viral proteins associated with RNA viruses are located in the cytoplasm. Immunolabeled dendrites, axon fibers, and cell somata showing small dots of dense viral antigen accumulation were detected in the parenchyma, mainly in the olfactory pathways, including the olfactory bulb (not illustrated), olfactory nuclei, olfactory tubercles, piriform cortex, and amygdala as well as the septum, ventral hippocampus, hippocampal fimbria, and polymorphic layer of the ventral dentate gyrus. Viral antigens were detected in both axons and dendrites, which frequently presented many abnormal varicosities sometimes associated with closely adjacent immunolabeled glial cells (not illustrated), suggesting a possible interaction between

![Figure 1. Cellular and neuroanatomical viral targets.](image)

Photomicrographs from Piry virus immunostained parasagittal sections of infected (A, C, and D) and uninfected (B) animals to illustrate cellular and targets areas at 8 dpi. Note the dendrites, axon fibers, and cell somata of neurons with altered cell appendages and microglia-like cells (arrows) in infected but not in control mice. Immunostaining includes the olfactory bulb, olfactory nuclei, olfactory tubercles, piriform cortex, septum (Sep), amygdala (Amyg. nu.), ventral hippocampus, hippocampal fimbria (F), polymorphic layer of the ventral dentate gyrus (DG), and CA3. ac: anterior comissure; H: hippocampus. Scale bars: low power: 1.0 mm; medium power: 250 μm; high power: 25 μm. doi:10.1371/journal.pone.0015597.g001
Response to Piry virus

Tomato lectin binds several major lymphocyte and microglial cell surface glycoproteins and is readily detected by a single two-step reaction, revealing the distribution of T cells and both quiescent and activated microglia [20,21]. Widespread microglial activation with an altered morphological phenotype was found at 8 dpi in the same areas where viral antigens were detected in both EE mice infected with Piry (IEPy) (Fig. 2Ai and Ci) and infected EE mice (EEPy) (Fig. 2Bi and Di), with a greater intensity in IE (Fig. 2A and C) than EE (Fig. 2B and D) animals. We have also found less viral immunolabeling in the brain sections from EEPy animals compared to IEPy (Fig. S1). In both IEPy and EEPy mice, at this time point, dorsal hippocampus showed less neuroinvasion (Fig. 2Ci and Di) than ventral hippocampus (Fig. 2Ai and Bi). Small rounded tomato lectin-positive cells that differed from the phagocytic phenotype of microglia were also detected around 8 dpi and were particularly prominent in the EEPy group (arrows in Fig. 2B and D). In the EEPy hippocampus, small cells were immunostained by anti-CD3, anti-CD8, and anti-CD43 (insert in Fig. 2B), suggesting that the EE had increased the occurrence of infiltrating T cells in the infected areas. After 20 dpi or later when the Piry virus had already been cleared from the brain parenchyma, infiltrated tomato lectin-positive cells was virtually absent. Uninfected EE control mice were devoid of rounded tomato lectin-positive cells and CD3 immunolabeled cells in the brain parenchyma.

Figure 2E represents average values of optical fractionator estimations of CA3-activated microglial cells in each environmental condition at 8, 20, and 40 dpi after histochemical staining with Lycopersicon esculentum agglutinin. Microglial estimations revealed a significant increase at 8 dpi in both IEPy and EEPy groups compared to the respective controls, but EEPy animals recovered to control levels at 40 dpi, whereas in EEPy animals, type I PNs were significantly reduced only at 8 dpi (one-way ANOVA, Bonferroni test \(p<0.05\)) but not in EEPy animals as compared to uninfected respective controls. However, type I and II PNs differed in their response; in IEPy mice, type I remained reduced until 20 dpi but recovered to control levels at 40 dpi, whereas in EEPy animals, type I PNs were significantly reduced only at 8 dpi (one-way ANOVA, Bonferroni test \(p<0.05\)). Type II PNs and totals (Type I plus Type II) were reduced in number only at 8 dpi in IEPy (one-way ANOVA, Bonferroni test \(p<0.05\)) but not in EEPy animals.

Two-way ANOVA revealed a significant influence of environmental conditions (\(F = 20.71; p<0.0001\)) and survival time (\(F = 38.73; p<0.0001\)) after infection on the total number of microglia in CA3, with a significant interaction between these variables (\(F = 4.18; p = 0.0138\)) whereas the number of type I PNs was influenced by the environment (\(F = 15.41; p = 0.0004\)) and survival time after infection (\(F = 3.12; p = 0.039\)) with a nonsignificant interaction between these variables (\(F = 0.11; p = 0.95\)). However, two-way ANOVA applied to the type II PNs revealed no significant differences between the experimental groups, suggesting that type I PNs are more sensitive to the viral infection and that the extracellular matrix regeneration is faster in EE than in IE animals.

Finally, the number of CA3 neurons did not differ between IEPy and EEPy mice or compared to their respective control groups at 20 dpi, regardless of environmental conditions (Table S4; IEcont: 35742±1920; IEPy: 3724±8480; EEcont: 35104±3438; EEPy: 33496±4825; mean ± s.d.; one-way ANOVA, \(p>0.05\)), indicating that microgliosis, PNs reduction, and behavioral changes were not associated with neuronal death.

The variance introduced by methodological procedures was, in most cases, less than 50% of the observed group variance, giving a ratio of CE2/CV2<0.5, where CE corresponds to the coefficient of error introduced by the stereological procedures and CV is the coefficient of variation [24]. In the cases of experimental groups that did not follow this rule, a CE2/CV2 ratio >0.5 was not detected.
indicative of a large variance introduced by stereological analysis. In this exception, both the CV and CE were low, and the general rule was neither meaningful nor practical to follow [24]. It has been previously demonstrated a direct correlation between microglial activation and extracellular matrix damage [25]. Table S5 list the coefficients of correlations between microglial numbers and different types of PNs estimations by the optical fractionator at different time windows. A significant inverse correlation was detected between the number of microglias and the number of type I PNs at 8 dpi ($p = 0.001$; coefficient of correlation $= -0.72$) and 20 dpi ($p = 0.008$; coefficient of correlation $= -0.63$) suggesting that the inflammatory response may contribute to the observed PNs damage during Piry viral encephalitis.

Figure 2. Viral neuroinvasion and microglial inflammatory response. Photomicrographs from IEPy (A, C, Ai, and Ci) and EEPy (B, D, Bi, and Di) parasagittal sections to illustrate the intensity of microglial activation and the presence of Piry virus antigens in the brain parenchyma. Note that the ventral CA3 (Ai, Bi), but not dorsal (Ci, Di), is intensely immunostained for Piry virus antigens and that morphologically activated microglia are more prominent in the ventral CA3 (A, B) as compared to the dorsal CA3 (C, D). Tomato lectin-positive small rounded cells other than activated microglia are indicated in EEPy (arrows). High-power pictures of immunolabeled CD-3–, CD-8–, and CD43-positive cells from the same region are illustrated in the insert (Bi). E: Total number and respective standar error bars (s.e.m.) of microglia estimations in CA3 at 8, 20, and 40 dpi and in control animals. Note the higher number of microglia in IEPy and that the numbers of activated microglia remained above control values, even after 40 dpi in IEPy, whereas EEPy returned to control levels at 20 dpi. Two-way ANOVA for environmental conditions ($F = 20.71; p < 0.0001$) and survival time ($F = 38.73; p < 0.0001$) with a significant interaction between these variables ($F = 4.18; p = 0.0138$). IEcont, EEpcont: impoverished and enriched environment control groups, IEPy,EEPy: impoverished and enriched environment infected groups. Scale bars: 25 μm (low power) and 10 μm (high power). doi:10.1371/journal.pone.0015597.g002
Behavioral changes

The first behavioral changes became apparent at 8 dpi, when IEPy animals burrowed significantly less food than IE controls and continued until 13 dpi, when burrowing activity recovered to control levels (Figure 5, top). No significant differences were detected in burrowing between EEPy and EE control animals (Figure 5, middle). Locomotor and exploratory activities assessed by open-field tests appeared altered later in the disease (20 dpi) and remained so at 40 dpi in IEPy animals, whereas no significant change was observed in EEPy mice (Figure 5, bottom). No significant differences were detected in the dark/light box or elevated plus maze tests.

Discussion

In the present report, as a model to study encephalitis outcomes in adult albino Swiss mice, we selected the Piry virus, a member of a group of RNA South American viruses [26,27], found in Brazil, that causes febrile disease in humans [28,29] and encephalitis in a neonate murine model [19,30,31]. In mice housed under IE or EE conditions, we induced viral encephalitis by intranasal inoculation of Piry virus–infected brain homogenate and correlated neuropathological features quantified using a stereologically based unbiased method with behavioral changes, comparing the outcomes with those of animals inoculated with uninfected brain homogenate.

Although a few reports describe application of stereological methods to quantify neuropathological features in correlation with behavioral changes to study encephalitis [32,33,34], none have investigated the effect of an EE on viral encephalitis. Indeed, only two studies have investigated the effect of an EE on brain infections using experimental models [35,36], but their approach to quantifying neuropathological changes was not based on unbiased stereological estimations. As a result, quantitative associations between neuropathological features and behavioral changes in murine models of encephalitis and the effect of EE have previously not been firmly established.

To quantify neuropathological changes, we applied the optical fractionator, an accurate method of quantification combining properties of an optical dissector and the fractionator that has been...
Microglial activation and neuronal death

The occurrence of neuronal death after encephalitis induced by rhabdoviruses remains controversial [41,42,43,44], but in general, when it is unequivocally detected, it seems to be a microglial-mediated event (e.g. [45,46]). To investigate possible losses of neuronal numbers in correlation with microglial activation after Piry virus encephalitis, we used NeuN as a selective immunohistochemical marker. The microglial host response was more intense and generalized in the brain parenchyma at 8 dpi compared to 20 or 40 dpi. Viral neuroinvasion mainly included the olfactory pathways, septum, amygdala, and ventral CA3 and polymorphic layer of the ventral dentate gyrus. There was an association between the intensity of viral antigen labeling in the parenchyma, a higher number of microglia, and a greater reduction of PNs, especially type I, without significant neuronal death. In addition, the intense immunolabeling of T lymphocytes in the brain was associated with environmental enrichment suggesting a higher mobilization of these cells to the brain parenchyma during Piry viral encephalitis. These findings suggest that microglia activation and extracellular matrix damage may be key factors in the pathogenesis of Piry encephalitis and that an EE differentially regulates microglial activation, increases T cell infiltration, preserves the extracellular matrix, and promotes faster virus clearance from the brain.

Neuroinvasion, the anti-inflammatory environment, and virus clearance

In the albino Swiss mouse model of viral encephalitis, microglial activation revealed by tomato lectin histochemistry occurred relatively early during disease progression (8 dpi) when the first behavioral changes became apparent. Tomato lectin also binds to monocytes and B and T infiltrating lymphocytes [21,22], revealing a conspicuous accumulation of lymphocytes in virus-infected areas. In the present report, these small rounded tomato lectin-positive cells, morphologically distinct from ameboid microglia, accumulated in infected areas in greater concentrations in animals housed under EE conditions when compared to those housed in impoverished conditions. Of importance, no apparent difference was found between uninfected animals in EE and IE conditions in terms of the distribution of T cells in the brain parenchyma. A significant number of CD3- and some CD8- and CD43-positive cells were found in the same regions (e.g., fimbria, septum) where tomato lectin-positive cells were detected. These results are compatible with previous data on VSV encephalitis [15,47].

Intranasal application of VSV induces acute encephalitis characterized by a pronounced myeloid and T cell infiltration with two distinct phagocytic populations regulating VSV encephalitis but not virus clearance [48]. VSV encephalitis is characterized by a pronounced infiltrate of myeloid cells (CD45+, CD11b+) and CD8+ T cells containing a subset specific to the immunodominant VSV nuclear protein epitope [15]. However, because ablation of peripheral macrophages does not impair VSV encephalitis or viral clearance from the brain but depletion of splenic marginal dendritic cells impairs this response and enhances morbidity/mortality [48], it is tempting to speculate that these dendritic cells may also be increased in EEpy as compared to IEPy animals. Another possibility is that the EE may induce NK cell mobilization of these cells to the brain parenchyma during Piry encephalitis or viral clearance from the brain but depletion of splenic marginal dendritic cells impairs this response and enhances morbidity/mortality [48], it is tempting to speculate that these dendritic cells may also be increased in EEpy as compared to IEPy animals. Another possibility is that the EE may induce NK cell activity previously described as absent in a VSV encephalitis murine model maintained in standard cages [15]. In line with these views, voluntary exercise such as that observed in an EE murine model maintained in standard cages [15]. In line with these views, voluntary exercise such as that observed in an EE murine model maintained in standard cages [15]. In line with these views, voluntary exercise such as that observed in an EE murine model maintained in standard cages [15]. In line with these views, voluntary exercise such as that observed in an EE murine model maintained in standard cages [15]. In line with these views, voluntary exercise such as that observed in an EE murine model maintained in standard cages [15]. In line with these views, voluntary exercise such as that observed in an EE murine model maintained in standard cages [15].
and ipsilateral reactive astrogliosis in a rat model of chronic inflammatory pain [56], induce beneficial astrogenesis to preserve the nigrostriatal system against 6-OHDA-induced toxicity [57], and alter the inflammatory response in the hippocampus of a transgenic mouse model of Alzheimer disease, TG2576 [58]. More recent work indicates that exercise training improves macrophage innate immune function in a beta(2)AR-dependent and -independent manner with lipopolysaccharide-stimulated nitric oxide and proinflammatory cytokine production in macrophages from trained mice being markedly higher than those from control mice [59].

In the present report, we expanded these observations, demonstrating that environmental conditions significantly affect disease outcomes in sublethal encephalitis. Because previous work had revealed that Piry virus neuroinvasion targets hippocampal fields [19] and that Wisteria floribunda histochemical staining conspicuously defines the architectonic limits of the hippocampal fields in adult mice [23], we chose to limit the quantitative neuropathological analysis to CA3. EE reduced the neuroinflammatory response and extracellular matrix damage in CA3. These findings are in agreement with those of previous reports suggesting that regular exercise provokes anti-inflammatory cytokines [60]. These previous data found an association between IL-6 and acute exercise, which was followed by stimulation of the production of anti-inflammatory cytokines and cytokine inhibitors, such as IL-1ra and IL-10, as well as an increase in IL-6 receptors during regular exercise. This favorable anti-inflammatory environment may explain the beneficial effects of exercise on acute neuroinflammation associated with the EE, as described in the present report.

As the disease resolved (20 and 40 dpi), the increased number of microglial cells in all infected animals and migratory CD3-positive cells in EE mice decreased and approached control levels after virus clearance. In line with these findings, the number of PNs increased in the infected animals in an inverse proportion to the activated microglial numbers in CA3, recovering the integrity of the extracellular matrix and normal behavior. These events were independent of the number of neurons in CA3 that remained unaltered during the disease course.

**Behavioral outcomes**

Morphologically activated microglia following Piry virus neuroinvasion were observed mainly in the olfactory pathways, septal region, amygdala, ventral CA3, and the polymorphic layer of the ventral dentate gyrus. Because septal damage mimics the effects of both dorsal and ventral hippocampal lesions [61], we selected ventral and dorsal hippocampal-dependent tasks to investigate behavioral changes [62,63,64,65]. Performances on all of these tasks are altered after hippocampal damage [65,66,67].

Burrowing changes started at 8 dpi and recovered to control levels at 13 dpi. Open-field tests presented significant differences between IEPI and IE control mice at 20 dpi and remained altered after 40 dpi. In contrast, animals housed under EE conditions had no significant differences in these tests. Assuming that the open field detected possible anxiety-like behavior associated with ventral hippocampus damage [64,65,68], that burrowing activity detects selective damage of the dorsal hippocampus [62,63,69], and that there was no apparent virus immunolabeling in the dorsal hippocampus, burrowing changes may be associated with septal damage [61].

In the murine model of VSV encephalitis, reactive astrogliosis and microglial activation occur relatively early in the disease [15,70]. As the disease progresses, these non-neuronal cells proliferate with an increasing effect on the extracellular matrix [71]. In the present report, microgliosis and a reduction in type I PNs in CA3 of IE mice were significantly correlated at 8 and 20 dpi, suggesting that the inflammatory response may be related to extracellular matrix damage. As soon as microglia activation was reduced during the disease recovery process, type I PNs started to recover up to control levels. Because the integrity of the extracellular matrix is important for long-term potentiation in the hippocampus [72], it may be possible that the observed type I perineuronal losses correlated, at least in part, with the transient behavioral changes observed with Piry virus encephalitis.

**Conclusion**

We report for the first time that an EE induces less intense behavioral changes, a lesser degree of microgliosis, a smaller reduction in the number of PNs, a higher degree of T cell infiltration, and faster virus clearance and disease resolution when compared to animals exposed to impoverished housing. We also demonstrated that nasal instillation of Piry-infected brain homogenate into adult albino Swiss mice induces (i) encephalitis with neuroinvasion, mainly of the olfactory pathways, septum, amygdala, and ventral hippocampus; and that (ii) the infection leads to an increase in CA3 microglial number and reduction of the PNs at 8 dpi when behavioral changes first appear, without changes in the number of neurons.

The mechanisms of neuronal protection that are activated during the faster clearance of the viruses from the brains of EE animals remain to be investigated. Detailed cellular and molecular analysis built on these observations, including characterization of the inflammatory cells mobilized to the brain parenchyma as well as viral neuroinvasion and clearance mechanisms, may delineate the pathophysiological basis of these events, improving our understanding of non-pharmacological treatment of neurological disorders.

**Materials and Methods**

All procedures were submitted to and approved by the institutional animal care committee of the Federal University of Paraí. We used 59 2-week-old albino Swiss mice obtained from the Animal Care Facility of Instituto Evandro Chagas and handled in accordance with the “Principles of Laboratory Animal Care” (NIH).

**Experimental groups and inoculation**

Suckling mice were intracerebrally infected with 10 μl of infected brain suspension, anesthetized, and perfused after becoming sick. The brains were histologically processed for immunohistochemistry procedures using specific anti-virus antibodies. Virus-containing brain homogenates were obtained as follows. First, 0.02 ml of each viral suspension was intracerebrally inoculated into newborn mice, and the animals were observed daily. Upon presenting with clinical signs, the animals were sacrificed and immediately stored at −70°C. Later, the brain tissue (0.2 g/animal) was macerated and mixed with 1.8 ml phosphate-buffered saline (PBS) containing 100 U/ml penicillin and 100 mg/ml streptomycin. The suspension was cleared by centrifugation at 10,000 x g for 15 min at 4°C. Virus titration was carried out by intracerebral inoculation of newborn mice with 0.02 ml of serial 10-fold dilutions of the viral suspensions in PBS, and LD50 values were calculated by the method of Reed and Muench. The Piry virus titers in the sample (LD50/0.02 ml) were 8.0 Log10. The choice of viral concentration obeyed the criterion of using a non-lethal dose that still induces a sublethal encephalitis.
in adult mice, achieved with a dilution of 1:100,000 of the sample titrated.

To confirm the presence of the virus in the brains of animals used to prepare infected brain homogenate, some of the animals were processed to obtain ultrathin sections for analysis with a Zeiss EM 900 transmission electron microscope, as described elsewhere [73]. In brief, samples obtained from neonate brains after perfusion and craniotomy were cut into small fragments and immersed for 2 h in a fixative solution containing 2.5% glutaraldehyde in 0.1 M phosphate buffer, pH 7.2, at room temperature. After primary fixation, brains were immersed in 0.1 M cacodylate buffer and post-fixed in a solution containing 1% osmium tetroxide, 0.8% potassium ferrocyanide, and 5 mM CaCl$_2$ at room temperature in the same buffer. Sections were en bloc stained with 2% uranyl acetate in 25% acetone, dehydrated in graded acetone concentrations, and embedded in EMbed-812 (Electron Microscopy Sciences, Fort Washington, PA, USA). Ultrathin sections were obtained with a Reichert/Leica Ultracut S (Electron Microscopy Sciences, Fort Washington, PA, USA) and stained with aqueous uranyl acetate and lead citrate before examination. Figure S2 shows a representative electronmicrograph of a neonate cortical region obtained by transmission electron microscopy to illustrate the typical bulb morphology of the Piry virus in the neonate brain used to prepare the infected brain homogenate.

Behavioral analysis

The mice were kept in an impovished environment (IE, $n = 31$) or enriched environment (EE, $n = 28$) for three months and then submitted to the following tests: open field, burrowing, dark/light box, and elevated plus maze. Enriched environmental conditions corresponded to plastic cages (32 cm x 39 cm x 16.5 cm) with chopped rice straw on the floor and equipped with road bridges, tunnels, running wheels, and toys made of plastic, wood, or metal with different forms and colors that were changed every week. The IE cages corresponded to plastic cages with the same dimensions and chopped rice straw on the floor but without equipment or toys. Each cage housed 12–15 mice. All mice had free access to water and food, and 12-h dark and light cycles were maintained. Tests occurred during the light cycle.

**Burrowing:** Two hours daily (from 09:00 to 11:00 h) for 3 consecutive days before inoculation and from post-inoculation days 2 to 35, all animals were placed in individual plastic cages (32 cm x 39 cm x 16.5 cm) containing a PVC tube (20 cm long, 7.2 cm diameter) filled with 250 g of normal diet food pellets. The open end was supported 3 cm above the floor. After the testing period, the remaining food in the cylinders was weighed and the mice returned to their collective cage [66].

**Open field:** The apparatus consisted of a gray PVC box (30 cm x 30 cm x 40 cm) with the floor divided into 10-cm squares. For 3 consecutive days before inoculation, each animal was placed in one corner and kept in the apparatus for 5 min. One meter above the open field, a video camera connected to a computer recorded each training session for later analysis by Any-Maze software (Stoelting). The following parameters were analyzed: distance traveled (m), mean speed (m/s), and immobility time (s). After each session, the open field was cleaned with 70% ethanol.

**Elevated plus maze:** The elevated plus maze consisted of two open (30 cm x 3 cm, no border) and two closed arms (30 cm x 5 cm, surrounded by a 15-cm wall) placed in opposite positions and connected by a central platform (5 cm x 5 cm). The apparatus was elevated 45 cm from the floor. Each animal was placed in the central platform facing one of the open arms and left there for 5 min. The test was performed over 2 consecutive days, and each animal had one session per day. All sessions were recorded and the following parameters analyzed by Any-Maze software (Stoelting): number of entries, time spent, and distance traveled in the open and closed arms. The program was set to define an arm entry when the center of the body of the animal entered the new area. For the sake of comparative analysis, the parameters were expressed as contrast values between the open and closed arms according to the following equation: $C = (c-o)/(c+o)$, where $C$ represents the contrast index and $c$ and $o$ correspond to parameters for the closed and open arms, respectively. The application of the contrast formula normalized the scale and allowed us to compare the anxiety-like behavior between groups with distinct patterns of locomotor activity, measuring possible differences more accurately.

**Dark/light box:** We adapted the dark/light box test from a previously published protocol [74]. The apparatus consisted of an open-topped rectangular box (45 cm x 27 cm x 30 cm high) divided into small (18 cm x 27 cm) and large (27 cm x 27 cm) areas with an open door (7.5 cm x 7.5 cm) located in the center of the partition at floor level. The small compartment was painted black and kept at a dim light level (0.38 cd/m$^2$), whereas the large compartment was painted white and brightly illuminated (36.4 cd/m$^2$). The test was performed in a quiet, dark room. All sessions were recorded with a webcam and the following parameters were analyzed by Any-Maze software (Stoelting): number of entries, time spent, and distance traveled in the light compartment. The program was set to define a compartment entry when the center of the body of the animal entered the new area. The mice were kept in this room at least 1 h before the test. To reduce any neophobic response to the test situation, the light/dark compartments were previously soiled by mice other than those used during the test. Mice were always tested in a soiled apparatus, and there was no cleaning between trials. Naïve mice were placed individually in the middle of the light area facing away from the opening. The images of the light compartment were recorded during a 5-min test.

For 6 consecutive days before inoculation, all mice were submitted to open field (days 1 to 3), LCE (days 4 and 5), and dark/light box (day 6) to obtain a baseline curve for these tests. After 20 days post-inoculation (dpi), 29 animals were submitted to the same tests, and the remaining mice ($n = 30$) were tested with the same protocol after 40 dpi.

After behavioral tests, all animals were anesthetized with intraperitoneal 2,2,2-tribromoethanol 1% (0.01 ml/g of body weight) and intra-nasally challenged with 5 μl viral suspension (10$^{-3}$ v/v in 100 U/ml penicillin, 100 μg/ml streptomycin) or normal brain homogenate as a control (10% v/v in 100 U/ml penicillin, 100 μg/ml streptomycin).

After recovering, animals were housed in enriched or standard plastic cages (32 cm x 39 cm x 16.5 cm) and maintained in the Instituto Evandro Chagas (Belém – PA) animal house, where they remained until the end of the experiments. All animals were kept in the care facility room at 23 ± 2°C with ad libitum access to food and water and with a 12-h light/dark cycle.

All mice were tested again (open field, burrowing, dark/light box, and elevated plus maze) after 20 dpi (IEPy, experimental, $n = 10$; IE control, $n = 7$; EEPy, $n = 5$; EE control, $n = 7$) or 40 dpi (IEPy, $n = 8$; IE control, $n = 6$; EEPy, $n = 11$; EE control, $n = 5$).

**Neuropathology**

When each animal reached the survival time of its group after behavioral tests, the mice were weighed and anesthetized with intraperitoneal 2,2,2-tribromoethanol (0.04 ml/g of body weight) and transcardially perfused with heparinized saline, followed by
4% paraformaldehyde in 0.1 M phosphate buffer (pH 7.2–7.4). Alternate series of sections (70 μm thickness) obtained using a Vibratome (Micron) were immunohistochemically labeled with polyclonal antibody for Piry virus antigens, monoclonal antibody for γ-glutamyl transpeptidase, and biotinylated lectin Wisteria floribunda to detect astrocytes, or monoclonal antibody for CD3- and CD8-positive T cells or histochemically reacted to detect the biotinylated lectin Lycopersicum esculentum (activated microglia) and Wisteria floribunda (PNs) supplied by Vector Laboratories (CA, USA). All chemicals used in this investigation were supplied by Sigma-Aldrich (Poole, UK) or Vector Laboratories (CA, USA), and the GFAP and Piry primary antibodies were from Chemicon (CA, USA) and Instituto Evandro Chagas (PA, Brazil), respectively.

Immunohistochemistry and histochemistry

To assess the distribution of Piry viral antigens, CD8 T cells, and astrocytes in the mouse brain at the different time points, immunohistochemistry was performed on all infected and in five control animals. Specific antibodies against Piry virus species were produced by the Unit of Arbovirus and Hemorrhagic Fevers at the Instituto Evandro Chagas, as described elsewhere [73,75]. In brief, free-floating sections were rinsed in 0.1 M phosphate buffer and placed in a solution of 0.2 M boric acid (pH 9.0) at 70°C for 1 h for antigen retrieval. After being rinsed in 0.1 M PBS with 5% Triton X-100, sections were incubated in a solution of methanol and 0.3% hydrogen peroxide for 10 min. After drying in PBS, the Mouse-on-Mouse (MOM) Blocking Kit (M.O.M. kit, Vector Laboratories, Burlingame, CA, USA) was used as follows: MOM IgG blocking for 1 h, primary antibody for 72 h (GFAP 1:800, Chemicon, CA, USA; anti-CD3 T lymphocytes 1:1000, MCA500G and anti-CD 43 MCA1096 1:200 AbD Serotec, Oxford, England, UK; anti-CD8 T lymphocytes 1:100, Cod 140063 eBioscience, San Diego, CA, USA; anti-Piry 1:20, Instituto Evandro Chagas, PA, Brazil), and MOM Biotinylated Anti-Mouse IgG Reagent for 12 h. Sections were washed in PBS and transferred to avidin-biotin-peroxidase complex (ABC) (Vector Laboratories) solution for 1 h, washed again before incubation in 0.2 M acetate buffer (pH 6.0) for 5 min, and revealed in GND solution (diaminobenzidine 0.6 mg/ml, ammonium nickel chloride 2.5 mg/ml, and glucose oxidase). All steps were carried out under gentle and constant agitation. All chemicals used in this investigation were supplied by Sigma-Aldrich (Poole, UK). As a negative control, normal horse serum was added to some slides instead of primary antibody for each antibody used as a cell marker and processed for immunohistochemistry as previously described.

Other sections were used to detect microglia activation and PNs by histochemistry with biotinylated Lycopersicum esculentum (tomato) lectin and biotinylated Wisteria floribunda lectin, respectively, according to manufacturers’ instructions with small adaptations. Briefly, sections were rinsed in PBS with 5% Triton X 100 for 20 min and transferred to a solution of methanol and 0.1% hydrogen peroxide for 10 min. After washing in PBS, the sections were incubated in the lectin solutions (Lycopersicum esculentum, 6 μg/ml; Wisteria floribunda, 9 μg/ml) overnight at 4°C, placed in ABC solution for 1 h, and revealed with GND solution following the same protocol previously described for immunohistochemistry. After this process, all sections reacted for Lycopersicum esculentum were counterstained in 0.5% cresyl violet.

Microscopy and optical fractionator

Details of the optical fractionator methodology are described under Stereological Procedures in the online supporting material (see Text S1). Tables S6–S8 present the stereological parameters and counting protocol and results for microglia and PNs in CA3.

Area and objects of interest: The limits of CA2/CA3 were defined by architectonic differences in the neuropil after Wisteria floribunda histochemistry, in which CA2 appears darker than CA3 (Figure S3). The PN counting procedure included two types of nets: type I corresponding to perisomatic and peridendritic nets, including secondary tertiary branches, and type II corresponding to perisomatic and faint primary dendrites (Figure S3).

We used histochemical reactions to reveal biotinylated Lycopersicum esculentum and Wisteria floribunda as markers of activated microglia and PNs. Detection of poly-N-acetyl lactosamine residues with biotinylated Lycopersicum esculentum (tomato lectin) also reveals T cell distribution and quiescent microglia [20,21]. Wisteria floribunda histochemistry selectively labels the n-acetyl-galactosamine B1 residues of glycoproteins, revealing PNs within the extracellular matrix and conspicuously defining the architectonic limits and layers of mice hippocampal fields CA3/CA2/CA1 [23]. Neurons were selectively labeled by immunohistochemistry to detect NeuN, a nuclear protein present in the vast majority of post-mitotic neuronal cells in vertebrates [76]. In the mouse hippocampus, where the area of interest for the present report is located, an extensive series of studies has already established NeuN as a selective neuronal marker [77,78].

Photomicrographic documentation and processing:

To obtain digital photomicrographs, we used a digital camera (Microfire, Optronics, CA, USA) coupled to a Nikon microscope (Optiphoto-2, NY, USA). Digital photomicrographs were processed using Adobe Photoshop 7.0.1 CS.2 software (San Jose, CA, USA) for scaling and adjusting the levels of brightness and contrast. For the figures, selected pictures were taken of sections from the animals in each experimental group with the total number of objects of interest nearest the mean value of each region of interest.

Statistical analyses

All groups were compared using parametric statistical analysis, one-way ANOVA, Bonferroni a priori test, or two-way ANOVA followed by Bonferroni post-tests, with differences between groups accepted as significant at a 95% confidence level (p<0.05).

Supporting Information

Text S1 Stereological Procedures

RTF

Figure S1 Cellular infection of Piry virus in EEPy and IEpy. Differential degree of Piry virus cellular infection in IE and EE infected subjects. Note less immunolabeled cells in EEPy (tomato lectin, red) compared to IEpy (primary antibody for each antibody used as a cell marker and processed for immunohistochemistry as previously described.

Figure S2 Electron micrograph of Piry virus in the cerebral cortex. Electron micrograph of Piry virus in the cerebral cortex of neonate albino Swiss mice used to prepare infected brain homogenate. Note the typical bullet morphology of this Rhabdovirus species (arrow heads). (TIF)

Figure S3 CA3 limits and perineuronal net types. Photomicrographs of histochemically reacted parasagittal sections of the architectonic limits of CA3 (low power) and types of perineuronal nets (high power). The CA3 pyramidal cell layer was outlined after histochemical reactions for Wisteria floribunda lectin (A) and immunohistochemistry for NeuN (not illustrated). Note that the Wisteria floribunda histochemical reaction labeled two types

Supporting Information

RTF

Figure S1 Cellular infection of Piry virus in EEPy and IEpy. Differential degree of Piry virus cellular infection in IE and EE infected subjects. Note less immunolabeled cells in EEPy as compared to IEpy (tomato lectin, red) compared to IEpy (primary antibody for each antibody used as a cell marker and processed for immunohistochemistry as previously described.

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of perineuronal nets, indicated as Type I and II. Arrows point to Type II and an asterisk to Type I perineuronal nets. Scale bars: low power 250 µm; high power 25 µm.

**Table S1** Microglial and perineuronal net estimations at 8 d post inoculation.

**Table S2** Microglial and perineuronal net estimations at 20 d post inoculation.

**Table S3** Microglial and perineuronal net estimations at 40 d post inoculation.

**Table S4** Neuronal estimations at 20 d post inoculation.

**Table S5** Correlations between microglial activation and extracellular matrix damage.

**Table S6** Stereological parameters for microglial estimations and counted markers.

**Table S7** Stereological parameters for perineuronal net estimations and counted markers.

**Table S8** Stereological parameters for neuronal estimations and counted markers.

**Author Contributions**
Conceived and designed the experiments: VHP CC PFCV JBTN JAPD CWPD. Performed the experiments: AAS RR NL NT AP ZS. Analyzed the data: JBTN CWPD VHP CC RR AAS. Contributed reagents/materials/analysis tools: PFCV JAPD CWPD CC VHP. Wrote the paper: CWPD CC VHP PFC JBTN.

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