INTRODUCTION

Current evidence suggests that chronic inflammatory diseases lead to neuroinflammation. Increased neuroinflammation can result in neurological impairment with deficits in cognition and motor function. For example, patients with diabetes, rheumatoid arthritis, obesity or chronic kidney disease can develop neurological deficits.[1-9] Inflammation and neuroinflammation are major contributing factors to cognitive and motor deficits in situations such as postoperative cognitive dysfunction, ageing and in some mental (e.g. schizophrenia) and neurodegenerative (e.g. Alzheimer’s disease) diseases.[10-16]

Patients suffering from chronic liver diseases (mainly cirrhosis and/or hepatitis) also show chronic inflammation which can lead to hepatic encephalopathy (HE): any alteration in cerebral function which is a consequence of previous liver failure.[17] There are two main types of liver diseases which induce HE: acute liver failure and chronic liver diseases. The effects and mechanisms underlying the cerebral alterations in acute and chronic liver failure are completely different.[17] This review will focus only on the mechanisms involved in HE in chronic liver diseases, mainly in liver cirrhosis.

Chronic liver diseases affect more than 5 million people in USA and a similar number in the European Union.[18] Patients with the chronic liver disease do not show neurological alterations at the beginning of the disease. However, with the progression of liver failure, most of these patients will suffer from some grade of HE. There are two main forms of HE in chronic liver disease: (1) minimal hepatic encephalopathy (MHE), in which the symptoms are not evident but can be unveiled using psychometric tests and (2) clinical or overt HE, in which the symptoms are evident. Once the symptoms are evident, clinical HE is graded in four stages according to the West Haven criteria.[19]

Approximately, 40-50% of cirrhotic patients present with MHE with mild cognitive impairment, attention deficits, psychomotor slowing, reduced mental processing speed and bimanual and visuomotor incoordination.[20-22] This incidence means that more
Inflammation and neuroinflammation in the neurological alterations in patients with chronic liver disease and MHE

Inflammation is a main contributor to the changes in cognitive and motor functions found in MHE patients suffering from chronic liver disease. For example, the serum levels of the pro-inflammatory cytokines IL-6 and IL-18 are increased in cirrhotic patients with MHE compared with cirrhotics without MHE. Moreover, the levels of IL-6 and IL-18 correlate with the grade of MHE evaluated using psychometric tests. Hyperammonemia plays a synergistic role with inflammation in the induction of the neurologic impairment (Figure 1). Hyperammonemia developed in cirrhotic patients by ingestion of an amino acid mixture induces neuropsychological impairment during inflammation, but not after its resolution. The contribution of inflammation and hyperammonemia to the neurological impairment in different pathological situations with different grades of inflammation and hyperammonemia was assessed by Felipo et al., who showed that the joint presence of inflammation and hyperammonemia, is enough to induce mild cognitive impairment, even in the absence of liver failure. The studies by Shawcross et al. and Felipo et al. support the idea that, as occurs in other chronic diseases, in chronic liver diseases, peripheral inflammation is involved in the induction of the cognitive and motor alterations associated to MHE and to overt HE.

Peripheral inflammation may lead to neuroinflammation by different mechanisms. The presence of neuroinflammation in patients with chronic liver diseases has not been thoroughly studied. One of the main methods to assess neuroinflammation is to assess by positron emission tomography (PET); the binding of ligands to the translocator protein (18 kDa) (TSPO), previously known as peripheral type benzodiazepine receptor. Cagnin et al. showed using PET techniques that cirrhotic patients with HE show increased binding of $^{[11C]}$(R)-PK11195 to TSPO in brain. Moreover, the patient with the most severe cognitive impairment had the highest increases in regional $^{[11C]}$(R)-PK11195 binding. The authors suggested that an altered glial cell state might be causally related to impaired brain functioning in HE. These data suggest the presence of neuroinflammation in patients with HE, correlating with the grade of cognitive impairment.

In support for the role of microglia, Dennis et al. performed immunohistochemistry studies and cytokine assays in fixed postmortem brain tissue from chronic alcoholics with cirrhosis and HE, alcoholics without HE and controls. About half of the patients with HE showed what they called "proliferative HE (pHE)" and the other half did not. The microglia in pHEs displayed an activated morphology with hypertrophied cell bodies and short, thickened processes. These patients also showed increased levels of IL-6. In contrast, the microglia in white matter regions of the non-proliferative HE cases were less activated and appeared dystrophic. The authors suggest that microglial proliferation may form part of an early neuroprotective response in HE that ultimately fails to halt the course of the disease because underlying etiological factors such as high cerebral ammonia and systemic inflammation remain. In support, Zemtsova et al. also analyzed markers of neuroinflammation in postmortem brain tissue from patients with cirrhosis that presented with and without HE and noncirrhotic
controls. They found an up-regulation of the microglial activation marker ionized calcium-binding adaptor molecule-1 (Iba-1) in the cerebral cortex from patients with cirrhosis who have HE but not from patients with cirrhosis who do not have HE. In this study, mRNA and protein expression of iNOS and cyclooxygenase (COX-2) or mRNA expression of pro-inflammatory cytokines and chemokine monocyte chemoattractive protein-1 was analyzed in cerebral cortex. They were not different in patients with liver cirrhosis and HE than in patients with cirrhosis who did not have HE or control patients without cirrhosis. This report shows increased microglial activation in patients with HE but did not find increased levels of pro-inflammatory markers.

Neuroinflammation has been also reported in patients with chronic hepatitis C (HC). However, in these patients a main contributor to neuroinflammation would be the presence of the virus into the brain. Postmortem studies of HC virus indicate that the viral replication might occur in the brain and microglia may be the locus for infection and subsequent neuroinflammatory activity. Bokemeyer et al. assessed the possible contribution of neuroinflammation to cognitive dysfunction in patients with HC who have mild liver disease. They performed an extensive neuropsychological examination and a magnetic resonance spectroscopy study in a group of patients with HC showing only mild liver disease but differing degrees of neuropsychiatric alterations. Choline, creatine and N-acetyl-aspartate and N-acetyl-aspartyl-glutamate concentrations in the basal ganglia were increased in the patients compared to controls. Fatigue correlated negatively with the metabolic changes. As the increase of choline, creatine and myo-inositol are usually interpreted to indicate glial activation and macrophage infiltration in chronic inflammation they conclude that HC virus infection may induce neuroinflammation and brain dysfunction. The negative correlation with fatigue suggests a cerebral compensatory process after infection. Grover et al. used two independent in vivo imaging techniques to assess the presence of neuroinflammation in patients with mild chronic HC and control subjects. Using magnetic resonance spectroscopy, they found that basal ganglia myo-inositol/creatine and choline/creatine ratios were significantly elevated in patients with chronic HC compared to controls. Using PET with a ligand for microglial/brain macrophage activation, Grover et al. found evidence of microglial activation, which positively correlated with HC viremia and altered cerebral metabolism. This suggests that the HC virus induces neuroinflammation, which would contribute to the cognitive alterations in patients with HC.

Although in patients with HC neuroinflammation seems to be caused mainly by the presence of the virus into the brain, the studies from Cagnin et al. and Dennis et al. and Zemtsova et al. suggest that neuroinflammation could also be present in patients with liver cirrhosis in the absence of virus infection.

**ROLE OF NEUROINFLAMMATION IN THE COGNITIVE AND MOTOR ALTERATIONS IN ANIMAL MODELS OF MHE**

Neuroinflammation would be a main contributor to the cognitive and motor alterations in minimal and clinical HE associated to chronic liver disease. This possibility is clearly supported by studies in animal models. Cauli et al. showed that rats with portacaval shunts (PCS), a main model of MHE, show neuroinflammation. Following PCS, rats showed increased levels of inflammatory markers such as IL-6 levels or COX activity in the brain. PCS rats show impaired ability to learn a Y maze task due to reduced function of the glutamate-nitric oxide (NO)-cGMP pathway in cerebellum. Cauli et al. showed that chronic treatment with the anti-inflammatory ibuprofen reduced neuroinflammation and restored the ability to learn the Y maze task because it restores the function of the glutamate-NO-cGMP pathway in cerebellum in vivo. This report showed for the first time that neuroinflammation is a main contributor to the cognitive impairment in MHE [Figure 1].

Neuroinflammation also contributes to other types of cognitive and motor alterations by altering other specific mechanisms modulating them. For example, neuroinflammation is a main contributor to hypokinesia in rats with MHE and chronic treatment with ibuprofen restored motor function. The mechanisms and brain areas involved in neuroinflammation-induced hypokinesia are different from those involved in impairment of learning in the Y maze. Hypokinesia in PCS rats is due to the increased level of extracellular glutamate in substantia nigra pars reticulata (SNr). Glutamate activates metabotropic glutamate receptor 1 (mGluR1) and induces an increase extracellular GABA in ventromedial thalamus (VMT). This, in turn, reduces extracellular glutamate in motor cortex, which reduces motor activity. Blocking mGluR1 in SNr with an antagonist normalizes GABA in VMT, glutamate in motor cortex and motor activity. A down-regulation of glutamate transporters contributes to the increased levels of extracellular glutamate in SNr. Chronic treatment of rats with MHE with ibuprofen normalizes the amount of glutamate transporters, reduces extracellular glutamate in SNr and normalizes motor activity.
activity.[43] The presence of neuroinflammation and its contribution to cognitive and motor alterations have also been demonstrated in the other main animal model of MHE: rats with bile duct ligation (BDL). Rodrigo et al.[44] showed that BDL rats have microglial activation, mainly in cerebellum, as demonstrated by immunohistochemistry. This is associated with increased levels of inflammatory markers [inducible nitric oxide synthase (iNOS), IL-1β, prostaglandin E2] and with cognitive impairment and hypokinesia. Similar to the previous studies, chronic treatment with ibuprofen reduced microglial activation and inflammatory markers and restored cognitive and motor functions in the BDL rats.[45] The above reports clearly support the idea that MHE in chronic liver failure is associated with neuroinflammation that affects different cerebral processes resulting in different types of cognitive and motor alterations [Figure 1].

**CHRONIC HYPERAMMONEMIA INDUCES ACTIVATION OF MICROGLIA AND NEUROINFLAMMATION**

Concerning the mechanisms by which chronic liver failure induces neuroinflammation, it seems that two main contributors would be involved: chronic hyperammonemia and peripheral inflammation.

In addition to PCS and BDL rats, a well-recognized rodent model of MHE are rats with “pure” hyperammonemia without liver failure induced by feeding an ammonium-containing diet.[41] As described above for rat model of MHE induced by PCS, rats with chronic hyperammonemia also show reduced function of the glutamate-NO-cGMP pathway in cerebellum[46] and reduced ability to learn the Y maze task,[47] which is restored when the function of the pathway is restored, for example, by treatment with sildenafil.[48] This suggests that hyperammonemia and neuroinflammation impair learning the ability by the same mechanism. This idea led Rodrigo et al.[45] to hypothesize that hyperammonemia would induce neuroinflammation in the brain, which would be responsible for impairment of the glutamate-NO-cGMP pathway and of cognitive function. To assess whether chronic hyperammonemia per se induces neuroinflammation, Rodrigo et al.[45] assessed in the brains of hyperammonemic rats without liver failure activation of microglia by immunohistochemistry and the levels of inflammatory markers [iNOS, IL-1β, and prostaglandin E2 (PGE2)]. Hyperammonemic rats show microglial activation [Figure 1], mainly in cerebellum, and increased levels of inducible iNOS, IL-1β, and PGE2 which are associated with impaired function of the glutamate-NO-cGMP pathway and with cognitive impairment and hypokinesia.

Chronic treatment with ibuprofen reduced microglial activation and inflammatory markers and restored the function of the glutamate-NO-cGMP pathway in cerebellum and cognitive and motor functions in hyperammonemic rats.[44] In summary, this indicates that: (1) chronic hyperammonemia per se induces neuroinflammation; (2) neuroinflammation mediates the effects of chronic hyperammonemia on the function of the glutamate-NO-cGMP pathway in cerebellum and on cognitive and motor functions in hyperammonemic rats. As neurological alterations are mainly due to altered glutamatergic and GABAergic neurotransmission,[48,49] this suggests that neuroinflammation-mediated alterations in neurotransmission are mainly responsible for changes in cognitive and motor function in MHE and clinical HE [Figure 1].

Hyperammonemia may also alter neurotransmission by other mechanisms. For example, Thrane et al.[50] have shown that ammonia, at high concentrations, alters astrocyte potassium buffering, increasing extracellular potassium concentration and over-activating the Na⁺-K⁺-2Cl⁻ co-transporter isoform 1 (NKCC1) in neurons. The consequent depolarization of the neuronal GABA reversal potential selectively impairs cortical inhibitory networks. This altered GABAergic neurotransmission may contribute to the neurological alterations in hyperammonemia. The alterations in glutamatergic and GABAergic neurotransmission in hyperammonemia and HE have been recently reviewed by Cauli et al.[48] In addition to the contribution of hyperammonemia, peripheral inflammation would also contribute to induction of neuroinflammation in chronic liver disease [Figure 1].

The mechanisms by which hyperammonemia induces microglial activation and neuroinflammation have been poorly studied for the moment. Zemtsova et al.[50] studied the effects of large ammonia concentrations on rat microglia in vitro and in vivo. In cultured rat microglia, ammonia-stimulated cell migration and induced oxidative stress and an up-regulation of the microglial activation marker ionized calcium-binding adaptor molecule-1 (Iba-1). Up-regulation of Iba-1 was also found in the cerebral cortex from acutely ammonia-intoxicated rats. However, ammonia had no effect on microglial glutamate release, prostaglandin synthesis, and messenger RNA (mRNA) levels of iNOS, COX-2, and IL-1α/β, tumor necrosis factor α (TNFα), or IL-6, whereas in cultured astrocytes ammonia induced the release of glutamate, prostaglandins, and increased levels of IL-1β mRNA. Although this study was performed using ammonia levels much higher than those that can be found in chronic liver disease in vivo, these data suggest that ammonia per se
may activate microglia. Additional factors would act synergistically with hyperammonemia to induce pro-inflammatory factors and neuroinflammation in chronic liver failure.

**THERAPEUTIC IMPLICATIONS**

The *in vivo* studies reported above in animal models show that neuroinflammation would be a key therapeutic target to improve the cognitive and motor alterations in MHE and overt HE. However, different approaches can be used to reduce neuroinflammation in chronic liver disease, but it is important to take into account the possible secondary effects of each treatment, which may depend on the type and grade of liver disease in each patient. For example, several studies show that ibuprofen ameliorates cognitive and motor function in rats with MHE.[40,43,45] However, ibuprofen is a non-steroidal anti-inflammatory drug (NSAID), and this type of anti-inflammatory is not recommended in cirrhotic patients because they may induce kidney damage and, in patients prone to portal hypertension (a common liver disease complication), the risk of gastrointestinal bleeding is intensified with NSAIDs.[51,52] Therefore, alternative therapeutic approaches are needed to reduce neuroinflammation that would present no secondary side-effects. As activation of microglia is a main contributor to neuroinflammation in animal models of HE, Agusti *et al.*[53] hypothesized that reducing directly microglial activation would be useful to improve cognitive and motor function in HE without inducing secondary effects in the kidney. Inhibiting the p38 MAP kinase antagonizes microglial activation[54] and inhibitors of p38 are being tested as therapeutic tools in different inflammatory and autoimmune diseases.[55] Taking this into account, Agusti *et al.*[50] assessed whether inhibiting p38 with SB239063 reduces neuroinflammation and improves cognitive and motor function in rats with MHE due to PCS without affecting the kidney. This group showed that treatment with SB239063 reduced microglial activation and inflammatory markers (PGE2, COX activity, iNOS, IL-1β, TNFα) in brain and completely restored learning ability, motor activity and coordination in PCS rats without inducing kidney damage. This suggests that inhibitors of p38 would improve the neurological status in HE without inducing secondary effects in the kidney. This type of inhibitors is being developed by different pharmaceutical companies to treat different chronic inflammatory diseases. It is likely that they can be developed further and following extensive pre-clinical and clinical evaluation become available to patients suffering from MHE.

Additional studies to identify and test new targets to reduce neuroinflammation in MHE without inducing secondary effects would be very useful to develop new therapeutic tools to reverse the cognitive and motor alterations in patients with MHE and clinical HE associated with chronic liver diseases.

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