Tau protein in cerebrospinal fluid (CSF): a blood–CSF barrier related evaluation in patients with various neurological diseases

Sigurd D. Süssmuth\textsuperscript{a}, Hansotto Reiber\textsuperscript{b}, Hayrettin Tumani\textsuperscript{a,}\textsuperscript{*}

\textsuperscript{a}Department of Neurology, University of Ulm, Oberer Eselsberg 45, 89091 Ulm, Germany
\textsuperscript{b}Neurochemisches Labor, University of Göttingen, Göttingen, Germany

Received 8 December 2000; received in revised form 16 January 2001; accepted 17 January 2001

Abstract

Tau protein (tau) is primarily localised in neurons, and after brain parenchymal damage its release into cerebrospinal fluid (CSF) is increased. The particular influences of blood–CSF barrier function and of disease topography on CSF tau levels have not been studied yet. CSF tau concentrations determined by enzyme-immunoassay in various neurological diseases (n = 61) were not dependent upon blood–CSF barrier dysfunction. Significant elevation of tau levels in patients with meningoencephalitis and cerebral hemorrhage indicates brain parenchymal damage. In contrast, tau levels remained normal in patients with bacterial meningitis if encephalitic complications did not occur. In patients with Guillain–Barré syndrome tau levels were low. Increased tau levels in active multiple sclerosis compared to clinically nonactive states indicate axonal pathology in active disease. © 2001 Elsevier Science Ireland Ltd. All rights reserved.

Keywords: Tau protein; Cerebrospinal fluid; Brain specific proteins; Blood–cerebrospinal fluid barrier function
MRI or CT, neurophysiological findings and CSF analysis. Analysis of normal controls were omitted in this study since the specific performance characteristics of the assay as well as age-dependent control values have been described in detail by several other studies [4,9,14,19] including a report from our laboratory [10].

CSF tau levels were determined using a sandwich ELISA based on the original method of Vandermeeren et al. [19] (INNOTEST hTAU, Innogenetics, Zwijndrecht, Belgium). The assay was performed according to the protocol supplied with the kit, and CSF tau concentrations of the samples were estimated from standard curves made for each assay.

In four CSF samples with GBS the tau concentration was below the detection limit of the kit (59 ng/l). In these cases we assumed a CSF tau concentration of 50 ng/l for calculation of the median. We compared our results with the tau ranges of normal individuals given in the protocol. Statistical evaluation was done by Wilcoxon–Mann–Whitney U-test.

Fig. 1 shows the tau protein concentration in CSF of patients with different diseases including GBS, neuroborreliosis, and spinal canal stenosis associated with blood–CSF barrier dysfunction (elevated $Q_{\text{Alb}}$) and without brain parenchymal damage. There was no correlation between $Q_{\text{Alb}}$ and the tau level indicating that tau level in CSF is not influenced by the blood–CSF barrier dysfunction. The tau concentration with a median of 149 ng/l (range 132–237 ng/l) in CSF of patients with spinal canal stenosis was within the reported normal range. In this disease group, age-dependent concentration changes were seen as described in the literature [4,9,14]. All five patients with GBS showed the lowest CSF tau concentrations (values < 59 and 106 ng/l), even in cases with moderate to severe blood–CSF barrier dysfunction. Tau concentrations in all patients with peripheral neuroborreliosis were within the normal range (median 99 ng/l), independent of duration of symptoms. In order to investigate whether CSF tau concentration changes over disease course, we examined the CSF tau level of two patients in a control lumbar puncture 11 and 28 days after the first one. In both cases, we found a slight increase of 23.5 and 47% of CSF tau which were still within the normal range.

CSF samples from patients with brain parenchymal disorders due to inflammatory or to vascular origin were analysed. As shown in Fig. 2, there were remarkable differences in the levels of tau in different inflammatory diseases of the central nervous system (CNS). If diagnostic lumbar puncture was done in the early stage of bacterial meningitis within the first 2 days, tau levels appeared normal (median 95 ng/l; range between 60 and 180 ng/l). All these patients received early antibiotic treatment and recovered completely without neurological deficits. One patient with a moderate CSF tau increase of 310 ng/l had an additional ischemia of the cerebellum 18 days prior to lumbar puncture. Another patient with a CSF tau concentration of 624 ng/l on the fourth day of symptoms had a meningoencephalitic complication. No correlation was found between tau and CSF cell counts or the CSF lactate level in these cases (not shown). In contrast to bacterial meningitis, patients with damage of the brain parenchyma as in viral encephalitis showed a clear
CSF tau elevation ($P < 0.05; U$-test). Increased tau values ranged from 327 and 548 ng/l (both Varicella zoster encephalitis) up to 15 653 ng/l (Herpes simplex encephalitis). The tau concentration showed no obvious correlation to the individual outcome.

As shown in Fig. 3, CSF tau levels in patients with parenchymal damage due to brain infarction or intracerebral bleeding (ICB) were increased. Tau levels reached significantly higher values in patients with ICB as compared to patients with brain infarctions ($P < 0.05, U$-test).

In MS, CSF tau levels ranged between 80 and 321 ng/l (Fig. 4). Ten out of 17 patients showed CSF tau levels above the normal range. Analysing CSF tau levels in the MS group by differentiating patients according to clinically active vs. inactive stages, we found a significant elevation in CSF tau in the first group (median 188 ng/l mean age, 41.1 ± 12.5, and 109 ng/l mean age, 50.5 ± 16.2, respectively; $P < 0.05, U$-test).

To our knowledge this is the first report on the relationship between CSF tau concentration and the blood–CSF barrier function. Analysing CSF tau in GBS, neuroborreliosis, and spinal canal stenosis, we found no evidence for a correlation between tau level and $Q_{AB}$. In contrast to CSF proteins originating mainly from meningeal and spinal cord structures [12,17], CSF proteins from brain parenchyma such as tau are not influenced by the blood–CSF barrier dysfunction.

Patients with spinal canal stenosis as well as patients with neuroborreliosis showed normal CSF tau levels, indicating that damage to the nerve roots does not lead to an increase of CSF tau levels. Our results could not confirm CSF tau elevation in GBS, as reported in a previous study with elevated tau in 5 out of 11 GBS patients [19]. The extremely low levels of CSF tau in our GBS patients, even in cases with severe blood–CSF barrier dysfunction, could be the result of increased elimination or consumption of tau in some way. Moreover, it is possible that CSF factors induce conformational changes of tau protein. Since Vandermeeren et al. [19] used the same assay for tau detection, these findings are somewhat surprising, but CSF tau levels could depend on the severity and stage of disease, co-morbidity and type of the therapy.

Previous studies also reported elevated CSF tau levels in different neurological diseases. However, in most studies it remained unclear whether an elevated CSF tau level is diagnostically useful or whether it is to be used only as a surrogate marker in such different diseases as stroke, multisystem atrophy, epilepsy, or different forms of encephalitis [4,14]. We have found increased CSF tau levels in brain parenchymal diseases independent of etiologic origin, except for patients with bacterial meningitis. This is consistent with the localisation of the pathology in the early stage of bacterial meningitis confined to the meninges. Since in all these patients early antibiotic intervention was performed and encephalitic complications were prevented, tau levels remained within the normal range. We therefore suggest that elevated CSF tau in bacterial meningitis may be a marker for brain parenchymal complications. Consequently, we found elevated tau levels in diseases with viral meningoencephalitis. Compared to patients with HSV encephalitis, patients with VZV encephalitis showed only slightly increased CSF tau levels. This might be explained by hemorrhagic effects in HSV encephalitis, since patients with intracerebral hemorrhage had the highest CSF tau levels.

In the present study we found increased tau levels in CSF...
of some patients with MS. Interestingly, higher tau concentrations were detected during acute relapses in contrast to patients with clinically non-active stages. Kapaki et al. [5] recently reported similar results with increased tau protein levels in MS, but they found significant tau elevations in both the relapsing-remitting and the progressive subtypes. Since it is known that in MS acute axonal injury and brain atrophy continuously go on even in clinically silent phases [15,16], it is possible that tau levels generally reflect the ongoing axonal damage. Since it is likely that human oligodendrocytes contain tau as well [8], it appears possible that CSF tau elevation in MS is an effect of axonal injury and/or oligodendrocyte damage.

In summary, the findings of this study demonstrate: (1) that CSF tau level is not dependent upon blood–CSF barrier dysfunction; (2) that elevation of CSF tau is specific for brain parenchymal damage; (3) that in patients with bacterial meningitis elevation of CSF tau indicates encephalitic complications; (4) that elevation of CSF tau level is associated with clinically active MS.