Avidity of antineurocytoskeletal antibodies in Alzheimer’s disease patients

Libuse Noskova, Lenka Fialova, Ales Bartos, Tomas Zima

Aims. To optimise the ELISA method for the avidity of IgG antibodies against neurofilament heavy chain (NfH) and to determine the levels and avidity of anti-NfH antibodies in patients with Alzheimer’s disease (AD) and a healthy control group.

Methods. Various dilutions of sera and concentrations of urea and sodium chloride as chaotropic reagents were tested in the process of the ELISA optimisation. The levels and avidity of anti-NfH antibodies were determined in 30 patients with Alzheimer’s disease and 30 age-matched cognitively normal elderly adults.

Results. Sera dilution 1:200 and urea as a chaotrope in a concentration 6 mol/L were chosen to be the most suitable for the avidity assay of anti-NfH antibodies by ELISA. The results showed no differences in either level or avidity of IgG anti-NfH antibodies between AD patients and cognitively normal persons. The levels of anti-NfH IgG antibodies inversely correlated with their avidities.

Conclusions. We optimised the ELISA method for the determination of anti-NfH antibody avidity determination which is suitable for research of anti-NfH antibody avidity in patients with neurological diseases associated with neurocytoskeletal defects. The determination of serum anti-NfH antibody avidity in AD patients seems to have limited diagnostic significance.

Key words: Alzheimer’s disease, antibodies, avidity, heavy chain of neurofilament, neurofilament

INTRODUCTION

Alzheimer’s disease (AD) is the most frequent neurodegenerative disorder. Various alterations in the neurocytoskeleton are associated with the neurodegenerative process. When the axon is damaged or neurodegeneration is initiated, the structures of the neuronal cytoskeleton normally localised inside in neurons are released from the neurons into the interstitial fluid, cerebrospinal fluid and bloodstream.

The cytoskeletal proteins in the extracellular space as a result of various pathological processes may interact with the cells of the immune system and induce an inflammatory or autoimmune response. The levels of autoantibodies against neurocytoskeletal autoantigens in the serum might be potential biomarkers for neurodegenerative diseases involving continuous neuronal death. Moreover, the study of autoantibodies is becoming important in association with intensive investigation and clinical trials testing the possibility of immunotherapy in AD.

In addition to the changes in antibody levels, their avidity is a substantial qualitative characteristic and could also be of clinical significance. The examination of antibody avidity in clinical practice is usually used for the differentiation of primary and chronic infection or for monitoring the effectiveness of vaccination.

Current knowledge about the avidity of autoantibodies is less complex than that about antibodies against exogenous antigens. High-avidity autoantibodies may be associated with autoimmune processes in the organism, but decreased avidity of antibodies against the β-amyloid protein in Alzheimer’s disease has also been described. However, information about the avidity of neuron-specific antibodies is sporadic so far, although it is a relevant issue of antibodies, which may determine their protective or injurious potency.

Recently, we found increased levels of intrathecally synthesised antibodies against NfH and tau proteins in patients with Alzheimer’s disease. Serum levels of anti-NfH IgG antibodies in AD have also been studied, but we are not aware of any study evaluating the avidity of these antibodies. Therefore we decided to continue with a more detailed study of anti-NfH IgG antibodies in serum by the determination of their avidity in the patients with dementia.

The aim of this study was to design and optimise ELISA for the determination of anti-NfH IgG antibody avidity. Various methods and procedures for the measurement of antibody avidity have been designed. One of
the methods suitable for clinical use is ELISA (Enzyme-Linked Immuno-Sorbent Assay), which is modified by the addition of substances with a chaotropic effect on the immune complexes formed during the ELISA procedure. These agents are necessary for the disruption of the immune complexes containing weakly bound antibodies. The lower the antibody avidity is, the more immune complexes are disrupted and the released antibodies are then washed away after incubation with the chaotropic agent.

Using our optimised ELISA, we evaluated the avidity of the anti-NfH IgG antibodies in the serum of patients with Alzheimer’s disease and an appropriate control group. We aimed to better characterise the humoral immune response against neurocytoskeletal components and to consider a possible diagnostic significance of anti-NfH IgG antibody avidity.

MATERIAL AND METHODS

Participants

We examined two groups of elderly persons. Serum samples were obtained from patients at the Department of Neurology, Charles University. The AD patients were recruited from the Memory Clinic. The diagnosis was based on objective evidence of a progressive decline in cognition, functional and detailed neuropsychological assessments, hippocampal atrophy seen on brain magnetic resonance imaging (or computer tomography in case of contraindications), temporo-parietal hypoperfusion seen using single photon emission computed tomography, or increased total or phosphorylated tau proteins and/or decreased β-amyloid concentrations in cerebrospinal fluid using cut-offs established in our previous studies. The diagnosis of dementia due to AD had been already established before blood sampling. They had mild form of dementia according to MMSE scores (Tab. 1). Many of them have already died after long-term clinical decline. Subjects in the control group (n=30; mean age 72 years) were cognitively normal elderly adults having normal Mini-Mental State Examination (MMSE) scores. They were age-matched with the AD patients.

The basic characteristics of the two groups are shown in Table 1. The AD patients and patients in the control group did not differ in age and other parameters correlates with epidemiological findings.

Patients included in the study signed the informed consent. The study was approved by the Ethics Committee of the Third Faculty of Medicine, Charles University.

Only selected samples were used for the optimisation of ELISA. The influence of pH was tested in five samples (control and AD patients’ samples). A comparison of chaotropic agents was performed in 20 samples (10 controls and 10 AD patients).

Methods

ELISA for anti-NfH IgG antibody level and avidity determination in the serum

Anti-NfH antibody IgG levels were determined by ELISA originally described by Silber et al. and later modified for the purpose of our previous studies. Two kinds of chaotropic agents – NaCl and urea – in various concentrations were tested for avidity method optimisation. Solutions at different concentrations were chosen for initial experiment concentrations based on previous works. Solutions of NaCl were used at concentrations 0.25 mol/L, 0.5 mol/L, 1.0 mol/L, 2.0 mol/L and 3.0 mol/L and urea at 2.0 mol/L, 4.0 mol/L, 6.0 mol/L and 8.0 mol/L. Since pH values change in the chaotropic agent solutions of different concentrations, we also tested the influence of pH on the avidity determination. Therefore, the avidity experiments were performed with the chaotropic agent solution both at the actual (original) pH and at pH adjusted to the uniform value of 7.4. Serum samples were analysed in serial dilution from 1:50 to 1:400.

Procedure of ELISA for the determination of anti-NfH IgG antibody levels and their avidity

The heavy neurofilament subunit isolated from a bovine spinal cord (purity is > 98%, declared by the manufacturer, Progen, Germany) was used as the antigen. An aliquot from the stock solution of the antigen was diluted in carbonate immobilisation buffer (10 mmol/L NaHCO₃, 0.15 mmol/L NaN₃, pH 9.6) to the concentration of 2.5 μg/mL. One half of the wells of 96-well plate type Maxisorp (Nunc, Denmark) was coated with 100 μL of diluted antigen and the other half of the wells was coated only with immobilisation buffer. The plate was incubated overnight at 4 °C.

After the binding of the antigen, the wells were washed (wash solution 0.1 M NaCl + Tween 20) and blocked with 1% BSA (bovine serum albumin) fraction V in PBS (phosphate buffered saline), pH 7.4. Then, the serum samples in appropriate dilution (in 1% BSA fr. V in PBS) were applied in an amount of 100 μL. Serum samples were incubated in the wells for 2 hours at room temperature.

After five washings of the wells, the chaotropic agents NaCl or urea in tested concentrations or PBS alone were applied to the appropriate wells in an amount of 100 μL/well. It was followed by incubation for 10 minutes at room temperature. The wells were washed (4×).

Data are expressed as average ±SD

<table>
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<tr>
<th></th>
<th>Controls</th>
<th>Alzheimer's disease patients</th>
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<td>Number of patients</td>
<td>30</td>
<td>30</td>
</tr>
<tr>
<td>Age at blood draw (years)</td>
<td>72 ± 5</td>
<td>74 ± 8</td>
</tr>
<tr>
<td>Gender</td>
<td>63% females</td>
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<tr>
<td>Education (years)</td>
<td>15 ± 3</td>
<td>13 ± 3</td>
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<tr>
<td>MMSE score (0-30 points)</td>
<td>29 ± 1</td>
<td>19 ± 6</td>
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MMSE - Mini-Mental State Examination score
100 μL/well of secondary antibody (conjugate) against the Fc region of human immunoglobulin IgG conjugated with peroxidase (Southern Biotech, USA, diluted 5000x in 1% BSA fr. V in PBS) was added to the wells in the next step and incubated for one hour at room temperature. After washings (4×), 100 μL of peroxidase substrate TMB (tetramethylbenzidine) with H2O2 (TestLine, Czech Republic) was added to the wells. The enzyme reaction was stopped by adding 2 mol/L H2SO4 (100 μL/well) after incubation for 15-20 min at room temperature in the dark. Absorbance (A) of the wells was measured at 450 nm against a reference wavelength of 620 nm using a Tecan Sunrise Plate Reader (Tecan, Austria) with the Magellan 6 program.

The absorbance of the blank sample (serum replaced with diluent) was subtracted from the absorbance value of each sample. For comparative purposes, the same pooled serum was used as the internal control in all analytical series of patients’ samples (n=60). The intra- and inter-assay of the ELISA method did not exceed 10%.

### Expression of anti-NfH IgG antibody concentration

Because no appropriate generally-used standards are available for anti-NfH IgG antibodies, a pooled serum prepared by mixing sera from patients with high levels of anti-NfH IgG antibodies (according their absorbance in the previous ELISA determination) served as a calibration standard for the determination of the arbitrary units (AU) of anti-NfH IgG antibody concentration. A calibration curve constructed from the absorbance of our standard diluted geometrically from 1:50 to 1:400 was run on every plate. Arbitrary units of concentrations were determined from the calibration curve created in the Magellan 6 program (Tecan, Austria).

### Expression of the avidity value

The avidity of antibodies is expressed as an avidity index (AI), which is defined as the ratio of the absorbance or AU value for the well with a chaotropic agent to the absorbance or AU value of the well without this agent. Usually, AI is expressed in the percentage. The high-avidity antibodies are commonly classified as those with an avidity index higher than 60% and the low-avidity ones as those with an avidity index lower than 40%. The avidity index values of the antibodies from 40% to 60% are usually marked as a “grey zone”, when the decision about the avidity is not clear.

### Statistical analysis

Non-parametric statistics were used because the data were not normally distributed. This statement was verified using a graphical evaluation and Shapiro-Wilk’s test. The median and interquartile range were used for patients and the control group characterisation. The correlation was tested using the Spearman correlation coefficient. The statistical evaluation of paired measurements was tested using the Wilcoxon matched pairs test. The Mann-Whitney test was used for comparisons between groups. The level of significance was set to 0.05. All statistical calculations were done in the Statistica program (StatSoft, Czech Republic).

### RESULTS

#### Expression of the anti-NfH IgG antibody levels and avidity

The concentrations of anti-NfH IgG antibodies can be expressed in the values of absorbance (450 nm) or in the AU obtained from a calibration curve described in the Methods section. Non-linear regression with an extrapolation factor of 2 was used for the construction of a calibration curve. An example of the calibration curve for the expression of anti-NfH IgG antibody concentration in arbitrary units is shown in Fig. 1.

Levels of anti-NfH IgG antibodies in the AU correlated with their expression in the absorbance (R=0.6; P<0.0001; n=60; Spearman correlation coefficient). Similarly, the avidity index calculated from absorbance values correlated with the calculation of the avidity index derived from the AU (R=0.976 for urea 2 mol/L, P<0.0001; R=0.959 for urea 6 mol/L, P<0.0001; Spearman correlation coefficient).

### Optimisation of the ELISA method for anti-NfH IgG antibody avidity determination

To optimise the method for avidity determination we focused on (1) the optimal dilution of serum samples; (2) to test the effect of chaotrophic solution pH; (3) to find the optimal concentration of chaotropic agents and further (4) to select the suitable chaotropic agent.

#### Serum dilution

The best dilutions of sera that corresponded with the linear part of the calibration curve appeared to be 1:200 and 1:400. However, the higher dilution 1:400 was optimal only for samples with elevated levels of antibodies. The absorbance values of sera diluted 1:400 were too low for medium or low levels of antibodies and the results...
were less precise. It was evident for both tested chaotropic agents (urea and NaCl) especially, when their higher concentration was used.

**pH of the chaotropic agent solutions**

Both chaotropic agent solutions - urea and NaCl - generated a pH gradient depending on the concentration of the agent. The mean change in pH values was 0.25 for NaCl solutions and 0.1 for urea solutions. The pH effect of the chaotropic agent solution for avidity determination was tested by analysing five samples (controls and AD patients, low, medium and high values of anti-NfH IgG antibodies). The avidity of each sample was determined using a chaotropic agent solution, in which the pH was adjusted to a uniform pH value (7.4) as well as with a chaotropic agent solution with original pH value arisen at preparation of the solution. The pH of the chaotropic agent solution in various concentrations did not significantly influence the determination of the anti-NfH IgG antibody avidity. Therefore, it was not necessary to modify the pH of the chaotropic agent solution in the following experiments.

**Concentrations of chaotropic agents**

It was crucial to find a concentration of the chaotropic agent that would clearly differentiate antibodies with high avidity from those with low avidity.

Low concentrations of NaCl (0.25 mol/L and 0.5 mol/L) showed minimal effect on the disruption of the immune complexes. Differentiation between high- or low-avidity antibodies was allowed by higher concentrations of NaCl (1 mol/L, 2 mol/L and 3 mol/L). These concentrations of NaCl were used for the further assays. After testing an additional four samples, the concentration of 1 mol/L and 3 mol/L NaCl were selected for further study because no significant differences between the 1 mol/L and 2 mol/L NaCl concentrations were seen.

Higher concentration of urea (6 mol/L and 8 mol/L) influenced the disruption of immune complexes by a similar intensity. Because of the increased risk of denaturation of immune complexes in the presence of higher concentration of urea, we chose urea concentrations of 6 mol/L for avidity determination. A significant difference between 4 mol/L and 6 mol/L urea concentrations was not found. Finally, the 2 mol/L and 6 mol/L urea concentrations were chosen for further analyses.

The result of the avidity assay depends on antibody levels\(^1\). The higher concentration of the chaotrope may be important for samples with high levels of antibodies, in which a lower concentration of the chaotrope could be insufficient for the disruption of a greater amount of immune complexes. Therefore, we performed the avidity determination at two different concentrations of chaotropic agents in the following avidity tests. Significant differences were seen between 1 mol/L and 3 mol/L NaCl (\(P=0.0001\), Wilcoxon matched paired test) and 2 mol/L and 6 mol/L urea (\(P<0.0001\), Wilcoxon matched paired test) - Fig. 2.

![Fig. 2. Comparison of NaCl and urea as chaotropic agents in the ELISA for the avidity determination of anti-NfH IgG antibodies. The box plots include median (horizontal line —), values from the 25th to the 75th percentiles (boxes); bars (whiskers) above and below the box indicate the range of non-outliers. Non-outliers are indicated by triangles. Outliers and extremes are shown by circles and asterisks.](image)

A. Levels of serum anti-NfH IgG antibodies analysed by ELISA method in presence of chaotropic agents (NaCl and urea) in different concentrations. The results obtained in absence of chaotropic agents (PBS) are also shown for comparison. The results are expressed in arbitrary units. A significant differences were seen between 1 mol/L and 3 mol/L of NaCl (\(P=0.0001\)), 2 mol/L and 6 mol/L urea (\(P<0.0001\)) or 3 mol/L NaCl and 6 mol/L urea (\(P=0.000892\)).

B. Avidity of serum anti-NfH IgG antibodies analysed by ELISA in the presence of chaotropic agents (NaCl and urea) in different concentrations. The results of avidity are expressed as an avidity index (%). Significant differences were seen between 1 mol/L and 3 mol/L NaCl (\(P=0.0001\)), 2 mol/L and 6 mol/L urea (\(P=0.0001\)) or 3 mol/L NaCl and 6 mol/L urea (\(P=0.0002\)).

AU – arbitrary unit; NfH – heavy subunit of neurofilaments; PBS – phosphate buffered saline; M – mol/L
Fig. 3. Comparison of serum levels and avidities of anti-NfH antibodies IgG in the group of Alzheimer’s disease patients (n=30) and in the control group (n=30).

The box plots include median (horizontal line —), values from the 25th to the 75th percentiles (boxes); bars (whiskers) above and below the box indicate the range of non-outliers. Non-outliers are indicated by triangles. Outliers and extremes are shown by circles and asterisks.

A. Levels of serum IgG anti-NfH antibodies without a significant difference between group of Alzheimer’s disease patients and the control group

B. Avidity of anti-NfH autoantibodies in serum of Alzheimer’s disease patients and the control group. The results of avidity are expressed as an avidity index (%). A significant difference between these two groups was not found (P>0.05).

AD – Alzheimer’s disease; AI – avidity index; AU – arbitrary unit; NfH – neurofilament heavy subunit

Fig. 4. Relationships between levels and avidities of anti-NfH IgG antibodies. Avidity was determined by ELISA using urea 6 mol/L as chaotropic agent.

A. The relationship between levels and avidities of anti-NfH antibodies IgG in the control group (n=30). A significant inverse correlation was found (R=-0.55, P=0.00147).

B. The relationship between levels and avidities of IgG anti-NfH antibodies in the group of the Alzheimer’s disease patients (n=30). A significant inverse correlation was found R=-0.409, P=0.025).

AI – avidity index; AU – arbitrary unit; NfH – neurofilament heavy subunit
Comparison of NaCl and urea as chaotropic agents

The ability of two chaotropic agents (NaCl and urea) to distinguish between high- and low-avidity anti-NfH IgG antibodies was tested. Lower (1 mol/L for NaCl or 2 mol/L for urea) and higher (3 mol/L for NaCl or 6 mol/L for urea) concentration of these agents were selected and twenty serum samples (10 controls and 10 Alzheimer’s disease patients) were analysed. The analyses were assessed in triplicates for the samples and in duplicates for calibration samples.

The influence of chaotropic agents on avidity determination was assessed according to the results both in arbitrary units and avidity indices. Different abilities of chaotropic agents to disrupt the immune complexes were observed. Levels of anti-NfH IgG antibodies (in AU) in the presence of different concentrations of chaotropic agents are shown in Fig. 2.

The values of avidity indices are similar when the 1 mol/L NaCl and 2 mol/L urea are used. A significant difference was observed between 3 mol/L NaCl and 6 mol/L urea (P=0.00016; Wilcoxon matched paired test). A reliable estimation of antibody avidity requires about 50% decrease in an antibody binding after treatment of immune complexes by chaotropic agents in the ELISA procedure. Thus, urea in the concentration of 6 mol/L resulted in the most effective disruption of the immune complexes. Urea was evaluated as a suitable chaotropic agent for the immunochemical (ELISA) determination of anti-NfH IgG antibody avidity (Fig. 2B). The optimised ELISA method using 2 mol/L and 6 mol/L urea as a chaotropic agent was used for the avidity determination in the serum samples for the clinical part of our study.

Avidity of anti-NfH IgG autoantibodies in serum of Alzheimer’s disease patients

Using the optimised ELISA, the avidity of the anti-NfH IgG antibodies was studied in sera of Alzheimer’s disease patients and controls.

We found no significant differences either in the levels or in the avidities of serum anti-NfH IgG antibodies between patients with Alzheimer’s disease and those in the control group (Fig. 3AB). Also there was no significant correlation between anti-NfH IgG antibodies or their avidity and the relevant sociodemographic characteristics such as age of participants or sex (Spearman correlation coefficient, n.s.).

The levels of anti-NfH IgG antibodies inversely correlated with their avidities. The relationship between levels of antibodies and their avidities was expressed more closely in the control group than that of the AD group. The correlation in the control group was significant for both concentrations of urea used for avidity determination (urea 2 mol/L: R=0.38, P=0.038; urea 6 mol/L: R=0.55, P=0.00147, Spearman correlation coefficient). On the contrary, the relationship between levels and avidities of anti-NfH IgG antibodies was only seen in the presence of 6 mol/L urea in the AD group (urea 2 mol/L: R=0.151, n.s.; urea 6 mol/L: R=0.409, P=0.025; Spearman correlation coefficient) – Fig. 4AB.

DISCUSSION

Avidity represents an important feature of antibodies, which significantly influences the formation of immune complexes and affectivity of antibodies. In this study we focused on the avidity of antibodies against NfH, which belong to the cytoskeletal structure specific for nervous tissue. We optimised the avidity assay for IgG antibodies against NfH by assessing a suitable dilution of sera and by selecting a chaotrope and its concentration; we then examined sera of patients with Alzheimer’s disease and control individuals.

Avidity (or functional affinity) is defined as the overall forces that hold together an antigen with a corresponding antibody. It is dependent on the affinity between antigen-binding sites of the antibody and epitopes of the antigen as well as valences of the antigen and the antibody. ELISA is suitable method for the determination of antibody avidity for clinical purposes. Since non-covalent interactions in the certain immune complex are characteristic for a specific antibody and appropriate antigen, it is necessary to test the optimal conditions of the ELISA avidity assay independently for individual antibodies and to select the suitable chaotropic agents and their concentration or suitable dilution of samples. Chaotropic agents such as urea, ammonium thiocyanate, guanidine hydrochloride, diethylamine or sodium chloride have already been used for the determination of antibody avidity. The optimisation of ELISA was carried out in the presence of anti-NfH IgG antibodies with NfH more effectively than tested concentrations of NaCl. The concentration of 6 mol/L urea dissociated about 45% immune complexes and met the requirement about a 50% decrease in an antibody binding after adding chaotropic agents to immune complexes in the ELISA procedure. The optimised ELISA method used 2 mol/L and 6 mol/L urea as a chaotropic agent was used for the avidity determination in the serum samples for the clinical part of our study.

Research over recent years has suggested the participation of humoral autoimmune processes involved in the pathogenesis of AD, which significantly influences the formation of immune complexes and affectivity of antibodies. In this study we focused on the avidity of antibodies against NfH, which belong to the cytoskeletal structure specific for nervous tissue. We optimised the avidity assay for IgG antibodies against NfH by assessing a suitable dilution of sera and by selecting a chaotrope and its concentration; we then examined sera of patients with Alzheimer’s disease and control individuals.

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Multiple autoantibodies against various autoantigens were investigated in the serum and/or CSF of AD patients. One group of autoantibodies included those against structures specific for nervous tissue, such as neurofilaments (NfL, NfH), tau proteins, glial fibrillary acidic proteins and others. It seems that some autoantibodies may act pathogenically while others could have protective effects. Animal experiments suggested an association between anti-NfH antibodies and some manifestation of AD. Rats immunised with the NfH of Torpedo cholinergic neurons developed specific antibodies, which accumulated
in the perikarya and neurites of neurons in the septum and hippocampus and in white matter tracts in the rats’ forebrains. The immunised rats showed a significant deficit in short-term memory and in a reversal of choice paradigm in a position discrimination test in contrast to long-term memory, which was not affected. Oron et al. found that cognitive deficit in rats with experimental autoimmune encephalitis after immunisation by the cholineretic NFH evolved more slowly than the decrease in the density of cholineretic neurons in the septum of rats and the accumulation of IgG in the same brain area.

The use of certain autoantibodies as diagnostic or prognostic markers is under investigation. Levels of these antibodies were studied in sera and/or CSF (ref. 27,28,46). We were mainly interested in the antibodies against autoantigens originating from the neurocytoskeleton in past. We found the differences in avidities of anti-NFM and anti-tau antibodies in serum and CSF in the patients with multiple sclerosis. In addition, differences in avidities of various anti-neurocytoskeletal antibodies were seen. These are not surprising findings. Totland et al. also observed various avidities of two onconeural antibodies Hu and Yo in patients with paraneoplastic neurological syndromes. Therefore, it is important to test the avidity of each autoantibody individually.

The pathogenic autoantibodies are mostly characterised by a higher avidity for autoantigens in the target tissues, but the occurrence of low-avidity autoantibodies in immune-mediated diseases has also been described. Jianping et al. investigated the avidity of anti-AB antibodies in AD and found that both levels and avidities of anti-AB antibodies were statistically lower in AD patients than in the age-matched control group. They assumed that avidity declination would have a great influence on the clearance of AB by the cells of the immune system. We assumed similar changes in avidity of serum anti-NFH antibodies. Unfortunately, the comparison of anti-NFH IgG antibody levels and avidities showed no differences between AD patients and control. However, the avidity of antibodies may change with the progression of the disease. The follow-up study of individual AD patients might find a certain trend in the avidity of anti-NFH IgG antibodies in the serum.

A statistically significant inverse relationship between avidities and levels of anti-NFH antibodies in both groups was found. This might reflect that high-avidity anti-NFH IgG antibodies are bound in the immune complexes with their antigens. Conversely, the low-avidity antibodies form immune complexes less effectively and they tend to reach higher levels as free molecules.

CONCLUSION

We optimised the ELISA method in order to find anti-NFH antibody avidity determination. This method has been used for the study of anti-NFH antibody avidity in serum of patients with Alzheimer’s disease, but it is also suitable for research of anti-NFH antibody avidity in patients with other neurological diseases associated with neurocytoskeletal defects. Neither levels nor avidity of IgG anti-NFH antibodies differed between AD patients and cognitively normal persons. In the future, we plan to investigate if the levels and avidity of antineurocytoskeletal antibodies will change during the follow-up study. In addition, we are going to determine separately, free anti-NFH IgG antibodies and those bound in immune complexes.

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Author contributions: LN: study design, laboratory analysis, data analysis and interpretation, writing of the manuscript; LF: study design, data analysis and interpretation, writing of the manuscript; AB: collection and analysis of clinical data, clinical evaluation, general corrections; TZ: supervising the study.

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REFERENCES


