Serum neurofilament light chain
No clear relation to cognition and neuropsychiatric symptoms in stable MS

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Abstract

Objective
To explore the hypothesis that serum neurofilament light chain (sNfL) indicative of neuroaxonal damage may improve precise disease profiling with regard to cognition and neuropsychiatric symptoms, we analyzed potential associations of sNfL levels with cognitive test scores, fatigue, depression, and anxiety.

Methods
Patients with relapsing-remitting and secondary progressive MS (SPMS) underwent an elaborated assessment including MRI, various cognitive tests, and patient-reported outcomes. We determined sNfL levels by single molecule array (Simoa) assay. Relationships between sNfL, cognition, neuropsychiatric symptoms, and demographical data were analyzed using correlations, group comparisons, and regressions.

Results
In 45 clinically stable patients with MS (Expanded Disability Status Scale = 2.73 ± 1.12, disease duration = 10.03 ± 7.49 years), 40.0% were cognitively impaired. Mean sNfL levels were 16.02 ± 10.39 pg/mL, with higher levels in the SPMS subgroup (p = 0.038). sNfL levels did reliably link neither with the investigated cognitive and affective parameters nor with fatigue levels. The only relationship found in a small subgroup of patients with SPMS (n = 7) with visuospatial learning (r = −0.950, p = 0.001) and memory (r = −0.813; p = 0.026) disappeared when further controlling for age, educational level, and sex.

Conclusions
In patients with stable MS at less advanced disease stages, sNfL did not convincingly relate to cognitive performance, fatigue, depression, or anxiety and thus may not serve as a surrogate biomarker for neuropsychological status in such populations.

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In patients with MS, symptoms such as cognitive impairment (CI), fatigue, and mood disturbances have a detrimental influence on working ability and quality of life, even at early disease stages. Neuroaxonal damage, leading toward the disruption of structural and functional neuronal networks, contributes to the development and progression of irreversible disability including cognitive deterioration in patients with MS. Widespread axonal damage and the resulting disconnection of neuronal circuits has also a prominent role in the pathogenesis of fatigue and affective symptoms such as depression. Yet, in standard clinical care, accurate and regular cognitive and neuropsychiatric assessment is often not incorporated despite its clinical importance due to limited time and financial resources. Disease activity and burden is instead approached by traditional measures such as the Expanded Disability Status Scale (EDSS), the Multiple Sclerosis Severity Score, or the Multiple Sclerosis Functional Composite, which do not sufficiently mirror cognitive and neuropsychiatric status. New biomarkers are therefore needed that easily and sensitively assess tissue damage at the axonal level and thereby might predict clinically relevant outcomes such as cognition, fatigue, and mood disturbances.

Neurofilament light chain (NFL) is a promising molecular surrogate marker that indicates neuroaxonal injury in various diseases because it is released into CSF and blood as a subunit of a cytoskeletal protein in case of neuroaxonal damage. In MS, CSF NFL levels correlate with classical relapses and disability. Recently, the single molecule array (Simoa) immunoassay technology with unprecedented analytical sensitivity enabled the detection of subtle NFL levels even in serum samples and successfully linked serum NFL (sNFL) to neuroaxonal damage and therapy effects in MS. Few studies though specifically explored NFL as a biomarker of cognitive performance and fatigue in MS. Importantly, most of them measured NFL levels in CSF, although studies on sNFL levels and cognition, fatigue, depression, or anxiety are rare.

Here, we investigated whether we could confirm previous results relating CSF NFL and cognitive performance in MS by sNFL testing. We additionally examined whether sNFL is related to fatigue, depression, and anxiety in our sample of patients with MS with mild clinical deficits.

### Methods

#### Patient population

We consecutively recruited patients with RRMS and SPMS from local ambulatory settings during October 2016–September 2018. All patients had a confirmed MS diagnosis according to the revised McDonald 2010 criteria, had an EDSS score of ≤5.0, were aged between 18 and 60 years, were fluent in German language, and were free of relapses for at least the past 60 days. Diagnoses and disease courses were documented in patients’ most recent medical records by their referring neurologists and were all reviewed and verified by the authors (O.A. and A.R.). Patients were required to present at least with mild CI, defined as a Symbol Digit Modalities Test (SDMT) z score of ≤−0.5 and >−3.0, which was confirmed during the neuropsychological assessment. Exclusion criteria covered any acute neurologic (other than MS), systemic inflammatory, oncologic, or psychiatric condition, severe visual or hearing impairment not otherwise corrected, or upper extremity deficits that could interfere with neuropsychological testing performance. The final sample consisted of 45 patients.

#### Neuropsychological assessment and procedures

Demographical and clinical information was sampled by a trained neuropsychologist. Table 1 displays the cognitive tests and self-report questionnaires comprised in subsequent neuropsychological testing. Two psychologists double-checked data transfer from paper record sheets into electronic databases to avoid data entry errors and to ensure data quality.

#### Neurologic assessment, MRI, and measurement of sNFL

A clinical visit for standard neurologic examinations, MRI scanning, and blood sampling all in 1 day was scheduled in the context of the neuropsychological assessment, and most participants were seen within 4 weeks. A certified neurologist of the Department of Neurology scored each participant on the EDSS according to established standards. In a 3T MRI system (Siemens Skyra, Erlangen, Germany), we acquired sagittal T1-weighted sequences (magnetization-prepared rapid acquisition with gradient echo [MPRAGE]) with 1 mm isotropic resolution (repetition time [TR] = 2,300 ms, echo time [TE] = 2.98 ms, field of view [FoV] = 256 × 256 mm, and 192 slices) and sagittal fluid–attenuated inversion recovery sequences with

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**Glossary**

BICAMS = Brief International Cognitive Assessment for MS; BVMT-R = Brief Visuospatial Memory Test—Revised; CBV = cortical brain volume; CI = cognitive impairment; CIS = clinically isolated syndrome; CV = coefficient of variation; DGV = deep gray matter volume; FoV = field of view; FSL = FMRI Software Library; MPRAGE = magnetization-prepared rapid acquisition with gradient echo; NBV = normalized brain volume; NFL = neurofilament light chain; SDMT = Symbol Digit Modalities Test; sNFL = serum NFL; T2-LL = T2 lesion load; TE = echo time; TR = repetition time.

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by digital immunoassay using the Simoa NF-light Advantage Kit (Quanterix Corporation, Lexington, MA) on our previously described setup. Preparation of samples including dilution steps was performed according to the manufacturer’s instructions, and NfL levels were determined in duplicates, using the corresponding standard curve. The intra-assay coefficient of variation (CV) was assessed by measuring a quality control serum, with a resulting CV below 5%. We accepted results of duplicates as valid if the CV of both measurements was below 20%, which was true for all samples.

**Statistical analysis**

Cognitive performances below the fifth percentile according to each test’s manual were identified as impaired. Global CI was defined as impairment in at least 2 cognitive domains. Performance in the Brief International Cognitive Assessment for MS (BICAMS) battery was considered impaired if at least 1 subtest score was below the fifth percentile.

We first examined the relationships of sNfL levels with demographical, clinical, imaging parameters, cognitive, and patient-reported outcomes by Pearson’s correlation coefficient or Spearman’s rank correlation. Whether sNfL levels differ along with categorical variables (sex and CI) was examined through group comparisons. Partial correlations were calculated to adjust for potential subtype effects. In a next step, 2 groups (low and high sNfL based on median split) were separated and compared on their performance in each test and questionnaire, as well as on global CI, and other demographical and clinical parameters using Student t tests or the Fisher exact test for parametric and using Mann-Whitney U tests for non-parametric variables. Group comparisons were also controlled for age, educational level, sex, and MS subtype using analyses of covariance. Finally, we conducted linear regression analyses with test and questionnaire scores, as well as the number of impaired cognitive tests and domains as the dependent and sNfL as the independent variable. Logistic regression models were analyzed likewise to predict impairment in each cognitive and affective domain including fatigue, impairment in BICAMS, and global CI. In all models, age, educational level, sex, and MS subtype were entered as covariates. In addition, outliers with standardized residuals > 3 or < -3 were excluded from the analyses. Despite being clearly aware that the subsample sizes of patients with RRMS vs SPMS in this work are not appropriate to draw general conclusions, we conducted separate correlational analyses as described previously in the 2 disease subtypes. We therefore aimed at exploring whether there are potential differences in the association of sNfL with demographical, clinical, cognitive, and neuropsychiatric parameters according to the MS phenotype.

Analyses were performed with IBM SPSS 25, with a significance threshold set at p = 0.05.

**Table 1 Neuropsychological assessment**

| Domain                          | Test/questionnaire                                      |
|---------------------------------|---------------------------------------------------------|
| Information processing speed    | SDMT*                                                   |
|                                 | TMT-A                                                   |
| Short-term memory and learning  | VLMT direct recall*                                     |
|                                 | BVMT-R direct recall*                                   |
|                                 | Digit span forward                                      |
|                                 | Corsi block forward                                     |
| Long-term memory                | VLMT delayed recall                                     |
|                                 | BVMT-R delayed recall                                   |
|                                 | BVMT-R recognition                                      |
| Working memory                  | Digit span backward                                     |
|                                 | Corsi block backward                                    |
| Verbal fluency                  | RWT phonetic condition                                  |
|                                 | RWT semantic condition                                  |
| Executive functions             | TMT-B                                                   |
|                                 | RWT switching condition                                 |
|                                 | DKEFS TAS*                                               |
| Subjective cognitive problems   | PDQ-20                                                   |
| Fatigue                         | FSMC                                                    |
| Anxiety                         | HADS A                                                   |
| Depression                      | HADS D                                                   |

Abbreviations: BVMT-R = Brief Visuospatial Memory Test—Revised; DKEFS TAS = Delis-Kaplan Executive Function System Tower Test Total Achievement Score; FSMC = Fatigue Scale for Motor and Cognitive Functions; HADS = Hospital Anxiety and Depression Scale; PDQ-20 = Perceived Deficit Questionnaire; RRMS = relapsing-remitting MS; RWT = Regensburger Verbal Fluency Test; SDMT = Symbol Digit Modalities Test; SPMS = secondary progressive MS; TMT = Trail Making Test; VLMT = Verbaler Lern- und Merkfaehigkeitstests.

* Components of Brief International Cognitive Assessment for MS.
* Missing data on DKEFS TAS for 12 patients with RRMS and 2 patients with SPMS.
All patients participated voluntarily in the study and provided written informed consent. The study was approved by the Ethics Committee of the Heinrich Heine University (study number: 5531R, registration-ID: 2016055083) and was conducted in accordance with the principles of the Declaration of Helsinki.

**Data availability**
Anonymized data will be made available by the corresponding author on reasonable request from any qualified investigator.

### Results

#### Patient and sNfL characteristics
Demographical and clinical characteristics of the final sample are shown in Table 2. Importantly, no participant experienced a relapse or changed immunomodulatory treatment within the past 2 months before blood sampling. The mean EDSS score was 2.73 (RRMS: mean = 2.62 [SD = 1.09], SPMS: mean = 3.36 [SD = 1.18]; t(43) = −1.63, p = 0.110, r = 0.242), while EDSS data of the total sample were distributed unimodally and without any outliers. T2-LL (RRMS: median =

### Table 2  Main demographical and clinical characteristics

| Variable                             | Total sample (N = 45) |
|--------------------------------------|-----------------------|
| Age, y, mean ± SD (median; range; IQR) | 45.63 ± 10.53 (47.61; 18.06–60.11; 16.49) |
| Sex, female, n (%)                   | 39 (86.7)             |
| Education, n (%)                     |                       |
| Low                                  | 4 (8.9)               |
| Middle                               | 9 (20.0)              |
| High                                 | 32 (71.1)             |
| Disease course, n (%)                |                       |
| RRMS                                 | 38 (84.4)             |
| SPMS                                 | 7 (15.6)              |
| Age at onset, y, mean ± SD (median; range; IQR) | 35.10 ± 9.87 (35.97; 14.94–53.19; 15.5) |
| Disease duration, y, mean ± SD (median; range; IQR) | 10.03 ± 7.49 (8.08; 0.51–28.27; 12.20) |
| EDSS, mean ± SD (median; range; IQR) | 2.73 ± 1.12 (3; 0–5; 1.5) |
| Immunotherapy classification, n (%)  |                       |
| None                                 | 7 (15.6)              |
| First line                           | 24 (53.3)             |
| Second line                          | 14 (31.1)             |
| Time since last relapse, mo, mean ± SD (median; range; IQR) | 37.91 ± 40.97 (21.84; 2.89–158.85; 41.33) |
| Time since last immunotherapy change, mo, mean ± SD (median; range; IQR) | 40.65 ± 47.86 (24.89; 1.86–220.06; 41.22) |
| T2-LL, cm³, mean ± SD (median; range; IQR) | 10.17 ± 10.18 (5.97; 0.99–36.43; 11.75) |
| NBV, cm³, mean ± SD (median; range; IQR) | 1,441.92 ± 69.41 (1,455.23; 1,256.01–1,554.36; 111.45) |
| CBV, cm³, mean ± SD (median; range; IQR) | 591.15 ± 37.16 (591.77; 517.37–687.93; 45.14) |
| DGV, cm³, mean ± SD (median; range; IQR) | 42.31 ± 6.29 (42.78; 24.98–53.02; 8.17) |
| sNfL, pg/mL, mean ± SD (median; range; IQR) | 16.02 ± 10.39 (13.8; 4.55–51.6; 11.98) |
| BICAMS impaired, n (%)               | 18 (40.0)             |
| Global CI, n (%)                     | 28 (62.2)             |

Abbreviations: BICAMS = Brief International Cognitive Assessment for MS; CBV = cortical brain volume; CI = cognitive impairment; DGV = deep gray matter volume; EDSS = Expanded Disability Status Scale; educational level low = ≤10 years, medium = 11 years, and high = ≥12 years; first-line medication = induction approach drug (moderate efficacy, high safety), incl. interferon-β (n = 4), glatiramer acetate (n = 6), dimethyl fumarate (n = 13), and cyclophosphamide (n = 1); IQR = interquartile range; NBV = normalized brain volume; RRMS = relapsing-remitting MS; second-line medication = escalation approach drug (high efficacy, more safety risks), incl. natalizumab (n = 5), fingolimod (n = 6), alemtuzumab (n = 1), rituximab (n = 1), and daclizumab (n = 1); sNfL = serum neurofilament light; SPMS = secondary progressive MS; T2-LL = lesion load.

* Missing information on time since last relapse for 1 patient with SPMS.

* Missing information on lesion load, normalized brain volume, and cortical brain volume for 1 patient with RRMS.
Association of sNfL levels with demographical and clinical parameters

When comparing patients with low sNfL values to patients with high sNfL values, groups differed with respect to DGV, meaning that patients with high sNfL values had lower DGV. Taking effect sizes into consideration, associations were also indicated for CBV, age, and disease duration (table 5). We observed no other notable difference between patients with low sNfL values compared to high sNfL values with respect to all other demographical and clinical variables.

Association of sNfL levels with cognition and neuropsychiatric parameters

In the RRMS sample, sNfL levels did correlate neither significantly with any of the investigated cognitive performance measures or self-report questionnaires on fatigue, depression, and anxiety nor with the number of impaired tests or impaired domains. With reference to effect sizes, sNfL is suggested to relate with SDMT performance albeit not reaching significance (see table 4 for detailed results). We did not find differences in sNfL levels related to CI in BICAMS (not impaired: median = 12.00 pg/mL, impaired: median = 12.40 pg/mL, U = 154.00, p = 0.529, r = −0.106) and in the overall test battery (not impaired: median = 10.40 pg/mL, impaired: median = 13.80 pg/mL, U = 140.00, p = 0.504, r = −0.112). The same applies to the subgroup of 7 patients with SPMS for BICAMS (not impaired: median = 19.50 pg/mL, impaired: median = 29.40 pg/mL, U = 2.00, p = 0.381, r = −0.609) and the overall test battery (not impaired: median = 18.40 pg/mL, impaired: median = 20.00 pg/mL, U = 5.00, p = 0.857, r = −0.134). We observed, however, negative correlations with the Brief Visuospatial Memory Test—Revised (BVMT-R) direct recall, BVMT-R delayed recall, and Delis-

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**Table 3** Correlations between sNfL levels and demographical and clinical parameters

|                     | sNfL RRMS (n = 38) |  | sNfL SPMS (n = 7) |  |
|---------------------|-------------------|---|-------------------|---|
|                     | r      | p Value | r      | p Value |
| Age, y              | 0.216  | 0.194   | −0.126 | 0.789   |
| Educational level   | −0.163 | 0.329   | −0.316 | 0.490   |
| Disease duration, y | 0.291  | 0.077   | 0.139  | 0.766   |
| Age at disease onset, y | 0.046 | 0.785   | −0.135 | 0.773   |
| EDSS                | 0.124  | 0.457   | −0.019 | 0.968   |
| immunomodulatory treatment classification | 0.109 | 0.516   | 0.144  | 0.758   |
| Time since last relapse, mo | −0.017 | 0.921   | 0.531  | 0.279   |
| Time since last change in DMT, mo | 0.084 | 0.631   | 0.802  | 0.407   |
| T2-LL, cm³          | 0.305  | 0.067   | 0.180  | 0.700   |
| CBV, cm³            | −0.229 | 0.172   | −0.229 | 0.622   |
| DGV, cm³            | −0.262 | 0.117   | −0.408 | 0.363   |
| sNfL                 | −0.191 | 0.250   | −0.485 | 0.270   |

Abbreviations: CBV = cortical brain volume; DGV = deep gray matter volume; EDSS = Expanded Disability Status Scale; NBV = normalized brain volume; RRMS = relapsing-remitting MS; sNfL = serum neurofilament light; SPMS = secondary progressive MS; T2-LL = lesion load.

Reported are Pearson’s correlation coefficients (combination with interval variable) or Spearman’s rank correlation (association with ordinal variable).
Kaplan Executive Function System Tower Test Total Achievement Score in this subtype (see table 4 for detailed statistics).

When adjusting correlations in the total sample for MS subtype, we still found a negative relation between sNfL levels and BVMT-R performance with a moderate effect size, although not significant (direct recall: \( r_p = -0.285, p = 0.060 \); delayed recall: \( r_p = -0.283, p = 0.062 \)). When adjusting correlations for further variables that potentially influence cognitive performance (age, educational level, sex, and subtype), correlation coefficients diminished and none of the examined relationships turned out significant.

In the total sample, patients with high sNfL levels performed worse on the BVMT-R delayed recall (and direct recall by tendency) than patients with low sNfL levels (table 5). Despite considerable effect sizes, group differences according to sNfL status were no longer detectable when controlling for variables potentially influencing cognitive performance (age, sex, educational level, and subtype).

### Table 4 Correlations between sNfL levels and neuropsychological parameters

| | RRMS (n = 38) | | SPMS (n = 7) | |
|---|---|---|---|---|
| Cognitive test scores | | | | |
| SDMT | -0.296 | 0.071 | 0.320 | 0.484 |
| VLMT direct recall | -0.106 | 0.526 | 0.023 | 0.962 |
| VLMT delayed recall | -0.126 | 0.452 | 0.113 | 0.809 |
| VLMT recognition | 0.072 | 0.668 | 0.122 | 0.794 |
| BVMT-R direct recall | -0.214 | 0.197 | -0.950 | 0.001** |
| BVMT-R delayed recall | -0.228 | 0.169 | -0.813 | 0.026* |
| BVMT-R recognition | 0.087 | 0.604 | -0.742 | 0.056 |
| TMT-A | 0.094 | 0.368 | -0.075 | 0.872 |
| TMT-B | 0.120 | 0.415 | -0.245 | 0.597 |
| Digit span forward | 0.150 | 0.969 | 0.247 | 0.593 |
| Digit span backward | -0.136 | 0.309 | 0.035 | 0.941 |
| Block span forward | -0.006 | 0.574 | 0.395 | 0.381 |
| Block span backward | -0.170 | 0.472 | 0.416 | 0.353 |
| RWT phonetic | 0.079 | 0.636 | 0.160 | 0.732 |
| RWT semantic | -0.036 | 0.830 | 0.155 | 0.740 |
| RWT switching | 0.045 | 0.789 | 0.087 | 0.853 |
| DKEFS TAS | -0.191 | 0.350 | -0.901 | 0.037* |
| No. of impaired tests | -0.044 | 0.793 | 0.327 | 0.474 |
| No. of impaired domains | -0.060 | 0.721 | 0.327 | 0.474 |

Questionnaires

| | | | | |
|---|---|---|---|---|
| FSMC total | -0.051 | 0.763 | -0.562 | 0.189 |
| HADS A | -0.185 | 0.267 | -0.526 | 0.225 |
| HADS D | 0.012 | 0.941 | -0.410 | 0.361 |
| PDQ-20 total | -0.044 | 0.791 | -0.499 | 0.254 |

Abbreviations: BVMT-R = Brief Visuospatial Memory Test—Revised; DKEFS TAS = Delis-Kaplan Executive Function System Tower Test Total Achievement Score; FSMC = Fatigue Scale for Motor and Cognitive Functions; HADS = Hospital Anxiety and Depression Scale; PDQ-20 = Perceived Deficit Questionnaire; RRMS = relapsing-remitting MS; RWT = Regensburger Verbal Fluency Test; sNfL = serum neurofilament light; SDMT = Symbol Digit Modalities Test; SPMS = secondary progressive MS; TMT = Trail Making Test; VLMT = Verbaler Lern- und Merkfaehigkeitstest.

Reported are Pearson's correlation coefficients. Interpretation of \( p \) values: *\( p < 0.05 \) and **\( p < 0.01 \).
Table 5  Demographics and neuropsychological test scores of patients with low and high sNfL levels (mean ± SD [range]; or percentages for CI frequencies)

| Variable                  | Low sNfL (n = 22) | High sNfL (n = 23) | p Value | r     |
|---------------------------|-------------------|--------------------|--------|-------|
| **Demographics (selected)** |                   |                    |        |       |
| Age, y                    | 43.37 ± 10.03 (26.74–57.73) | 47.80 ± 10.77 (18.06–60.11) | 0.069 | −0.270 |
| Disease duration, y       | 7.71 ± 5.75 (0.51–20.86) | 12.25 ± 8.37 (0.79–28.27) | 0.084 | −0.257 |
| EDSS                      | 2.70 ± 0.97 (1–4) | 2.76 ± 1.27 (0–5) | 0.817 | −0.035 |
| T2-LL, cm³                | 7.67 ± 8.44 (0.99–35.21) | 12.66 ± 11.31 (1.21–36.43) | 0.127 | −0.230 |
| NBV, cm³                  | 1,458.05 ± 72.51 (1,256.01–1,554.36) | 1,425.78 ± 63.74 (1,303.04–1,503.66) | 0.124 | 0.235 |
| CBV, cm³                  | 43.62 ± 6.88 (24.98–53.02) | 41.06 ± 5.52 (28.83–51.74) | 0.048* | −0.294 |
| **Cognitive test scores** |                   |                    |        |       |
| SDMT                      | 44.68 ± 8.22 (24–58) | 41.74 ± 8.76 (26–61) | 0.252 | 0.174 |
| VLMT direct recall        | 56.50 ± 10.80 (27–73) | 51.43 ± 11.49 (23–70) | 0.135 | 0.226 |
| VLMT delayed recall       | 13.18 ± 2.13 (0–15) | 12.65 ± 2.39 (8–15) | 0.326 | −0.146 |
| BVMT-R direct recall      | 24.27 ± 5.88 (12–34) | 20.61 ± 7.88 (2–34) | 0.085 | 0.259 |
| BVMT-R delayed recall     | 9.73 ± 2.88 (2–12) | 8.13 ± 3.21 (0–12) | 0.029* | −0.326 |
| BVMT-R recognition        | 5.45 ± 1.34 (0–6) | 5.09 ± 2.04 (1–6) | 0.849 | 0.028 |
| TMT-A                     | 37.75 ± 15.21 (18.65–90.74) | 42.60 ± 14.20 (25.08–82.24) | 0.200 | −0.191 |
| TMT-B                     | 84.30 ± 30.71 (33.53–168.94) | 98.49 ± 50.50 (40.29–256.78) | 0.467 | −0.108 |
| Digit span forward        | 7.05 ± 2.01 (3–11) | 7.57 ± 1.67 (4–10) | 0.278 | −0.162 |
| Digit span backward       | 7.18 ± 1.97 (2–10) | 6.39 ± 1.70 (3–9) | 0.127 | −0.228 |
| Block span forward        | 8.36 ± 1.62 (5–11) | 8.48 ± 1.62 (6–12) | 0.835 | −0.031 |
| Block span backward       | 7.59 ± 1.97 (4–11) | 7.74 ± 2.12 (2–11) | 0.730 | −0.051 |
| RWT phonetic              | 19.32 ± 5.68 (5–28) | 19.17 ± 9.10 (4–37) | 0.937 | −0.012 |
| RWT semantic              | 31.59 ± 7.62 (14–41) | 28.91 ± 9.74 (12–42) | 0.488 | −0.103 |
| RWT switching             | 20.82 ± 4.00 (11–29) | 20.61 ± 4.65 (12–28) | 0.964 | −0.007 |
| DKEFS TAS                 | 19.36 ± 3.37 (13–25) | 17.35 ± 3.84 (12–23) | 0.175 | −0.202 |
| No. of impaired tests     | 2.68 ± 2.98 (0–9) | 3.00 ± 2.63 (0–11) | 0.363 | −0.136 |
| No. of impaired domains   | 1.91 ± 1.82 (0–6) | 2.22 ± 1.45 (0.5) | 0.312 | −0.151 |
| BICAMS CI                 | 36.36% | 43.48% | 0.763 | 0.073 |
| Global CI                 | 54.55% | 69.57% | 0.365 | 0.155 |

**Questionnaires**

|               | Low sNfL (n = 22) | High sNfL (n = 23) | p Value | r     |
|---------------|-------------------|--------------------|--------|-------|
| FSMC total    | 72.32 ± 12.76 (50–97) | 70.23 ± 14.47 (36–87) | 0.829 | −0.032 |
| HADS A        | 7.86 ± 3.58 (1–15) | 6.48 ± 3.67 (2–16) | 0.154 | −0.213 |
| HADS D        | 5.82 ± 4.50 (1–15) | 5.78 ± 4.35 (0–15) | 0.945 | −0.010 |
| PDQ-20 total  | 31.27 ± 12.94 (5–53) | 31.74 ± 12.16 (10–57) | 0.946 | −0.010 |

Abbreviations: BICAMS = Brief International Cognitive Assessment for MS; BVMT-R = Brief Visuospatial Memory Test—Revised; CBV = cortical brain volume; CI = cognitive impairment; DGV = deep gray matter volume; DKEFS TAS = Delis-Kaplan Executive Function System Tower Test Total Achievement Score; EDSS = Expanded Disability Status Scale; FSMC = Fatigue Scale for Motor and Cognitive Functions; HADS = Hospital Anxiety and Depression Scale; NBV = normalized brain volume; PDQ-20 = Perceived Deficit Questionnaire; RWT = Regensburger Verbal Fluency Test; sNfL = serum neurofilament light; SDMT = Symbol Digit Modalities Test; T2-LL = lesion load; TMT = Trail Making Test; VLMT = Verbaler Lern- und Merkfähigkeitstest. Reported are results of Student t tests, Mann-Whitney U tests, and the Fisher exact test. Interpretation of p values: *p < 0.05; r = correlation coefficient as effect size measure.
Finally, in the regression analyses, sNfL did not predict cognitive performance, fatigue, depression, or anxiety levels (neither the continuous performance scores in linear models nor categorical impairment in logistic models) (figure).

**Discussion**

In this study, we investigated whether and how sNfL levels are associated with cognitive and neuropsychiatric status in patients with MS.

In the examined group of patients with RRMS and SPMS, sNfL levels were comparable with previous studies using the same methodology, supporting the reliability of our results and of the applied assay analysis method. The observed prevalence rate of CI of 40% according to BICAMS and 62.2% when considering all assessed domains in this study also matches with former investigations. Notably, these percentages might have been overestimated because of the inclusion criteria of an SDMT scoring of at least $z \leq -0.5$. Mean performance scores of other tests (for instance of Verbaler Lern- und Merkfaehigkeitstest and BVMT-R) were average compared with normative values, albeit our sample was heterogeneously distributed considering demographical variables (such as age and educational level) to resemble patient populations in real-life settings. Patients with SPMS moreover did not differ from patients with RRMS in any of the cognitive test scores (details not reported), altogether suggesting that this sample’s cognitive functions were only mildly to moderately affected. The same can be noted for physical disability, as patients’ mean EDSS score was less than 3 (with maximum values not greater than 5) and did not differ between RRMS and SPMS. It can be presumed that only patients with relatively preserved clinical functioning, feeling capable of bearing a comprehensive testing, and making their way autonomously to and between the test centers registered for the study. Our MRI data compared with data from studies that particularly examined patients at higher functioning levels (i.e., patients with minimal physical disability or without CI) finally support the assumption that our sample has not been exposed to high neuronal disease activity on average.

Regarding demographical and clinical variables, we found sNfL to relate by tendency with age, disease duration, and imaging measures of lesion load and brain atrophy (T2-LL, CBV, and DGV). This agrees with the idea of sNfL potentially indicating axonal damage and neuronal degeneration in...
accordance with other numerous studies,\textsuperscript{15} although effect sizes were rather small.

Regarding cognitive performance and neuropsychiatric measures, sNfL was only related to visuospatial learning and memory. This seems to be mainly driven by patients with SPMS because correlational analyses in separate MS subtypes were significant only in this subgroup. These results align with a recent study that also examined BICAMS (with NfL from CSF) and observed the only domain-specific, tentative relation between NfL and BVMT-R performance. Besides, the authors did not find an association of BICAMS test performance within the subgroup of patients with RRMS as opposed to progressive patients. They argued that this might be traced to greater gray matter degeneration, expressed in higher NfL levels, in the progressive subtype.\textsuperscript{19} Visuospatial learning and memory were indeed defined within 1 cognitive cluster along with information processing speed, both being frequently and severely affected in the course of the disease and thus constituting sensitive domains for cognitive screening, particularly in SPMS.\textsuperscript{33,34} Other studies correspondingly observed information processing speed to be linked with NfL derived from plasma,\textsuperscript{35} CSF,\textsuperscript{17,18} and serum.\textsuperscript{36} Despite approaching a medium effect size, this association did not reach significance in our sample though. Jakimovski et al.\textsuperscript{36} for instance specifically measured sNfL with the same technique as in this study. They found sNfL to correlate with information processing speed in a considerably mixed sample of patients with relapsing and progressive MS, but not in patients with clinically isolated syndrome (CIS).

The results within the subsample of patients with SPMS in our study should be interpreted with caution because of the very small group size that potentially has biased statistical analyses and effect size estimations. In parallel, the overall sample size entails risk for lacking power in trying to detect associations between sNfL and demographical, clinical, cognitive, or neuropsychiatric parameters (i.e., type II error). We also need to emphasize that, given the exploratory nature of our study, the reported results are not corrected for multiple testing. In case of p-level adjustments, however, the observed associations would largely lose statistical significance. When controlling for factors that could additionally influence cognitive functioning (such as age, educational level, and sex), observed relationships likewise disappeared and sNfL did not predict any of the assessed cognitive and neuropsychiatric parameters, indicating that other parameters explain more variance of the outcome.

Taken together, we did not explore a clear link between sNfL and any cognitive test score, fatigue, depression, or anxiety levels. From a methodological perspective, these results can barely be attributed to an overly narrow selection of assessment scales and tools. We applied an extensive neuropsychological battery, not only covering various cognitive tests but also fatigue and affective self-report scales, assessing largely every neuropsychological domain NfL could potentially be associated with. Of interest, there are also other investigations that did not find CSF or sNfL to relate with cognitive performances\textsuperscript{20,22} or with baseline fatigue.\textsuperscript{21} Of note, they all examined cohorts consisting of patients with CIS or patients with recently diagnosed RRMS suggestive of low disease activity, in contrast to the studies mentioned above showing positive results. In this context, it should be highlighted that NfL was seen as a measure of acute inflammation, given its transitory increase during relapses.\textsuperscript{37} Most of our patients though were currently treated with DMTs known to effectively reduce disease progression and blood NfL levels.\textsuperscript{35,38} We found cognitive performance (especially in the domains of attention and information processing, as well as verbal and visuospatial learning and memory) instead to be correlated with MRI variables, with much stronger effect sizes than relations between sNfL and cognition or between sNfL and MRI (table e-1, links. lww.com/NXI/A311). Measures such as lesion load and atrophy of brain structures reflect accumulated damage over time and therefore might constitute better predictors of functional outcomes, which are assessed in the absence of acute disease activity. A recent study further indicated that NfL concentrations derived from CSF better mirror MS-related neuronal damage and degeneration than sNfL measures.\textsuperscript{39} Of note, the authors investigated a cohort with a similar size as compared to our population. As Håkansson and colleagues state “... levels of brain-derived markers in the blood still constitute a proxy for the levels in [the] CSF where CNS pathology may be better represented, at least in smaller cohorts....”

Considering all discussed aspects together, our results suggest that for populations with rather modest clinical manifestations and no acute disease activity, NfL measured from serum does not seem to be a surrogate biomarker for cognitive performance and neuropsychiatric symptoms. With the perspective of being applied in clinical contexts, the sensitivity of sNfL as a single measure for such complex functional outcomes particularly within small samples outside of large scientific trials is questioned and needs to be further examined in future studies.

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| Melanie Filser, MSc   | Cogito Center for Applied Neuropsychology and Neurological Research, Dusseldorf, Germany | Data acquisition and management, statistical analysis and interpretation, and critical revision and approval of the final version of the manuscript |
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Appendix (continued)

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| Philipp Albrecht, MD | Department of Neurology, Medical Faculty, University Dusseldorf, Germany | Data acquisition, and critical revision and approval of the final version of the manuscript                                                   |
| Markus Otto, MD | Department of Neurology, University Hospital Ulm, Germany | sNfL data analysis, and critical revision and approval of the final version of the manuscript                                            |
| Hayrettin Tumani, MD | Department of Neurology, University Hospital Ulm, Germany; Dietenborn, Ulm, Germany | sNfL data analysis, and critical revision and approval of the final version of the manuscript                                            |
| Iris-Katharina Penner, PhD | Department of Neurology, Medical Faculty, University Dusseldorf; Cogito Center for Applied Neuropsychology and Neuropsychological Research, Dusseldorf, Germany | Literature research, study design and conceptualization, acquisition of funding, data analysis and interpretation, manuscript drafting, and critical revision and approval of the final version of the manuscript |

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