CSF neurofilament light chain testing as an aid to determine treatment strategies in MS

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Abstract

Objective
To evaluate the use of CSF neurofilament light chain (NfL) measurements in clinical practice as well as their effect on treatment strategies and outcomes in patients with MS.

Methods
This was an observational cohort study of patients with MS who had a CSF NfL measurement between December 2015 and July 2018 as part of their routine clinical care. Treatment strategies were classified as “No Treatment/No Escalation” (no treatment or no escalation of treatment) or “Treatment/Escalation” (first-line injectable/oral disease-modifying therapies (DMTs), highly active DMTs, or treatment escalation). Change in Expanded Disability Status Scale (EDSS) scores was evaluated after 1-year follow-up.

Results
Of 203 patients with MS, 117 (58%) had relapsing-remitting MS. Disease activity was most frequently indicated by elevated CSF NfL (n = 85), followed by clinical (n = 81) and MRI activity (n = 65). CSF NfL measurements were independently associated with clinical (p = 0.02) and MRI activity (p < 0.001). Of those with elevated CSF NfL as the only evidence of disease activity (n = 22), 77% had progressive MS (PMS). In patients with PMS, 17 (20%) had elevated CSF NfL as the sole indicator of disease activity. Elevated CSF NfL resulted more frequently in Treatment/Escalation than normal CSF NfL (p < 0.001). Median EDSS change at follow-up was similar between patients receiving No Treatment/No Escalation and Treatment/Escalation decisions (p = 0.81).

Conclusions
CSF NfL measurements informed treatment strategies, alongside clinical and MRI measures. CSF NfL levels were the only indicator of disease activity in a subset of patients, which was more pronounced in patients with PMS. Elevated CSF NfL was associated with more Treatment/Escalation strategies, which had an impact on EDSS outcomes at 1 year.

*These authors contributed equally to the manuscript.


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The availability of new and more effective disease-modifying therapies (DMTs) has increased the complexity of MS management. Moreover, the clinical and pathologic heterogeneity throughout the disease course of MS poses major challenges for treatment decisions. Treatment strategies in MS are largely based on clinical activity (including relapses) and MRI findings (new or enlarging T2 lesions and/or gadolinium [Gd]-enhancing lesions). However, it is not possible to capture the full extent of disease activity with these measures, and the correlation between MRI measures and clinical disability remains limited. As a result, treatment strategies based solely on these activity markers may fail to deliver the best possible long-term outcomes.

Personalization of MS management is a key goal for all MS practices. There is, thus, an unmet need for additional biomarkers that enable neurologists to further stratify treatment strategies and improve outcomes for individual patients. As elevated CSF neurofilament light chain (NfL) measurements can indicate ongoing inflammation and neuroaxonal degeneration in MS, measurement of CSF NfL may represent an additional tool to assist in the treatment decision-making process. CSF NfL measurements have been shown to correlate with clinical and radiologic disease activity and predict disability progression. Moreover, reduction of CSF NfL measurements also indicates response to treatment. However, the utility of CSF NfL measurement in day-to-day clinical practice remains largely unexplored. Specifically, we do not know what form this will take over and above that of clinical and MRI activity and whether using CSF NfL in the treatment decision-making process has any impact on outcomes.

At our center, CSF NfL testing has been provided to assist treatment decision making since December 2015. In this cohort study, we aimed to (1) characterize the distribution of disease activity as measured by CSF NfL, clinical activity, and MRI activity, (2) evaluate the influence of CSF NfL measurements on treatment strategies, (3) evaluate the impact of CSF NfL-based treatment strategies on disability outcomes, and (4) evaluate the impact of the CSF NfL on our clinical practice following its introduction.

**Methods**

**Study design and participants**

This was an observational cohort study based at Barts Health NHS Trust, London, United Kingdom. Patients with either relapsing-remitting MS (RRMS) or progressive MS (PMS) who underwent CSF NfL measurements between December 2015 and July 2018 were identified from our institutional database. Inclusion criteria were age ≥18 years, having a treatment decision that took into account CSF NfL measurements, clinical and MRI assessments, and having an Expanded Disability Status Scale (EDSS) score obtained at least 1 year after the treatment decision.

**Glossary**

<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Definition</th>
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<tbody>
<tr>
<td>DMT</td>
<td>Disease-modifying therapy; EDSS = Expanded Disability Status Scale; Gd = gadolinium; IQR = interquartile range; NEDA = no evident disease activity; NfL = neurofilament light chain; PMS = progressive MS; RRMS = relapsing-remitting MS.</td>
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</table>
CSF NfL measurements (pg/mL) were calculated using a standard curve according to the manufacturer’s instructions. The detection limit of the ELISA was 33 pg/mL. Intra- and interassay coefficients of variation were below 10%. All NfL analyses were performed in duplicate. CSF NfL measurements were categorized as normal or elevated according to age-related reference values defined by the manufacturer. These reference ranges have been established in 50 volunteers that had no apparent neurologic disease based on interviews by a research nurse and MRI. The healthy subjects were divided into 3 age groups, and the reference levels were defined as median NfL level ± 2 SDs. In patients aged <30 years (n = 17), the cutoff was 290 pg/mL (median 186.4 pg/mL, 2 SD 100), in patients aged between 30 and 39 years (n = 15) 380 pg/mL (median 288.4 pg/mL, 2 SD 94.5), and in patients aged between 40 and 59 years (n = 18) 830 pg/mL (median 490.6 pg/mL, 2 SD 340).

**Treatment strategies**

The Barts MS center has implemented a local strategy of treating to a target of no evident disease activity (NEDA). Although we used CSF NfL measurements for NEDA assessment in MS, there was no specific algorithm for DMT selection based on NfL testing at our center. The decision on which DMT to prescribe was at the discretion of each MS consultant. Treatment strategies were classified as follows based on NHS England treatment algorithm and local prescribing policies: (1) “no treatment” when patients were not started on any DMT, (2) “no escalation of treatment” when previously treated patients continued on the same DMT, (3) “first-line injectable and oral DMTs” when treatment naive patients received beta-interferons, glatiramer acetate, teriflunomide, or dimethyl fumarate, (4) “highly active DMTs” when naive patients received high efficacy oral ( fingolimod), subcutaneous cladribine or infusion therapies (natalizumab, ocrelizumab, or alemtuzumab), and (5) “treatment escalation” when previously treated patients according to (2) and (4) were escalated to more effective DMTs. We defined treatment strategies (1) and (2) as “No Treatment/No Escalation” and (3), (4) and (5) as “Treatment/Escalation.”

**Statistical analysis**

Categorical variables were described as frequency and percentages and continuous and ordinal variables by median and interquartile range (IQR). CSF NfL levels were tested for normality using the Shapiro-Wilk test. As they were not normally distributed, pairwise comparisons were conducted with the Kruskal-Wallis test. Frequencies of grouped treatment strategies per year were compared using the χ² test. Two-way analysis of variance was performed to model CSF NfL measurements as a function of MRI and clinical activity taking into account covariates that were significantly associated with CSF NfL measurements. We checked for significant interactions between MRI and clinical activity and performed a sensitivity analysis in which extreme NfL values were replaced by the mean ± 3 SDs. A nominal significance threshold (p = 0.05) was applied, and all tests were 2 sided. All analyses were performed using the statistical package R v3.6.1.

**Data availability**

All data included in these analyses will be shared as anonymized data via request from any qualified investigator.

**Results**

**Patient characteristics**

A total of 203 patients with MS were included in the study, with a median age of 44 years (IQR 33–52 years), 123 (61%) were female and 117 (58%) had RRMS. The median EDSS score was 3 (IQR 1.5–6), and the median disease duration was 6 years (IQR 2–13 years). At baseline, 169 (83%) patients were not treated with any DMT (table 1).

**The distribution of disease activity as measured by CSF NfL, clinical activity, and MRI activity**

There was no evidence of clinical or MRI activity or elevated CSF NfL in 64 patients, whereas in 139 patients, at least 1 category signaled active disease (figure 1). Among those with disease activity, all 3 parameters were present in 21 patients (15%), while the frequency of disease activity was in the order of elevated CSF NfL (n = 85) > clinical activity (n = 81) > MRI activity (n = 65). In those without clinical activity, disease activity was still demonstrated in 39 patients through elevated CSF NfL and in 36 patients having an active MRI, with 17 patients displaying both (figure 1). CSF NfL was associated with MRI activity (p < 0.01) and with Gd-enhancing lesions (p < 0.001) (figure 2, A and B). CSF NfL was also associated with clinical activity (p < 0.001) and with relapses (p < 0.01) (figure 2, C and D).

When integrating both MRI and clinical variables in the same model, we demonstrated that NfL measurements were independently associated with clinical (b = 391.49, p = 0.02) and MRI activity (b = 766.31, p < 0.001) (figure 2E). No statistically significant interaction between MRI activity and clinical activity was observed in the model (p = 0.06). These results survived a sensitivity analysis (clinical activity: b = 252.96, p = 0.03; MRI activity: b = 566.94, p < 0.001). CSF NfL measurements were significantly higher in men than in women (p < 0.001) and were also associated with current treatment status (p < 0.01), which were both controlled for in the regression analysis.

**The influence of CSF NfL measurements on treatment strategies**

In 22 patients with MS (11%), the only evidence of disease activity used in the treatment decision-making process was their elevated CSF NfL measurement (“CSF NfL only” patients). The majority (77%) of these had PMS compared with 42% in the “MRI activity only” and 48% in the “clinical activity only” subgroup (table 1). CSF NfL only patients had a median disease duration of 8 years (IQR 4–10.8 years) and a median
Table 1 Baseline characteristics of the study cohort by disease activity statusa

<table>
<thead>
<tr>
<th>Characteristicb</th>
<th>Patients with MS (N = 203)</th>
<th>“NfL only” patients with MS (n = 22)</th>
<th>“MRI activity only” patients with MS (n = 19)</th>
<th>“Clinical activity only” patients with MS (n = 27)</th>
<th>“NfL and clinical activity” patients with MS (n = 17)</th>
<th>“NfL and MRI activity” patients with MS (n = 17)</th>
<th>“Clinical and MRI activity” patients with MS (n = 8)</th>
<th>“NfL, MRI activity, and clinical activity” patients with MS (n = 21)</th>
<th>“No activity” patients with MS (n = 64)</th>
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<tbody>
<tr>
<td>Sex, no. (%)</td>
<td></td>
<td></td>
<td></td>
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<td></td>
</tr>
<tr>
<td>Female</td>
<td>123 (60.6)</td>
<td>6 (27.3)</td>
<td>13 (68.4)</td>
<td>19 (70.4)</td>
<td>16 (64)</td>
<td>10 (58.8)</td>
<td>6 (75)</td>
<td>11 (52.4)</td>
<td>42 (65.6)</td>
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<tr>
<td>Male</td>
<td>80 (39.4)</td>
<td>16 (72.7)</td>
<td>6 (31.6)</td>
<td>8 (29.6)</td>
<td>9 (36)</td>
<td>7 (41.2)</td>
<td>2 (25)</td>
<td>10 (47.6)</td>
<td>22 (34.4)</td>
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<td>Age, median (IQR)</td>
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<tr>
<td>RRMS</td>
<td>44 (33–52)</td>
<td>46 (38.3–57)</td>
<td>46 (39–53)</td>
<td>49 (44.5–59)</td>
<td>33 (29.5–45.5)</td>
<td>35 (27–49.5)</td>
<td>38.5 (33.5–44.5)</td>
<td>33 (28–41)</td>
<td>47 (40–54)</td>
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<td>PMS</td>
<td>117 (57.6)</td>
<td>5 (22.7)</td>
<td>11 (57.9)</td>
<td>14 (51.9)</td>
<td>19 (76)</td>
<td>12 (70.6)</td>
<td>7 (87.5)</td>
<td>17 (81)</td>
<td>32 (50)</td>
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<tr>
<td>Disease duration, median (IQR)</td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>RRMS</td>
<td>6 (2–13)</td>
<td>8 (4–10.8)</td>
<td>8 (2.25–11)</td>
<td>9 (2–16.5)</td>
<td>4 (1–7)</td>
<td>5 (1–14)</td>
<td>3.5 (2.5–4.5)</td>
<td>2 (1–10)</td>
<td>9 (4–18)</td>
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<td>Baseline EDSS score, median (IQR)</td>
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<tr>
<td>RRMS</td>
<td>3 (1.5–6)</td>
<td>6 (4–6.5)</td>
<td>3 (1.5–6)</td>
<td>3 (2–5.3)</td>
<td>3 (1.5–5.3)</td>
<td>2 (1–4.3)</td>
<td>1.3 (1–1.5)</td>
<td>2 (1–3.5)</td>
<td>4 (2–6.5)</td>
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<td>DMTs, no. (%)</td>
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<tr>
<td>None</td>
<td>169 (83.3)</td>
<td>21 (95.5)</td>
<td>17 (89.4)</td>
<td>23 (85.2)</td>
<td>24 (96)</td>
<td>15 (88.2)</td>
<td>7 (87.5)</td>
<td>17 (81)</td>
<td>45 (70.3)</td>
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<td>—</td>
<td>—</td>
<td>—</td>
<td>1 (1.6)</td>
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<tr>
<td>Beta-interferon</td>
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<td>—</td>
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<td>—</td>
<td>—</td>
<td>1 (1.6)</td>
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<tr>
<td>Cladribine (subcutaneous)</td>
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<td>6 (3.0)</td>
<td>1 (4.5)</td>
<td>1 (5.3)</td>
<td>2 (7.4)</td>
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<td>—</td>
<td>1 (4.8)</td>
<td>1 (1.6)</td>
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<tr>
<td>Dimethyl fumarate</td>
<td>8 (4.0)</td>
<td>—</td>
<td>—</td>
<td>1 (3.7)</td>
<td>—</td>
<td>1 (5.9)</td>
<td>1 (12.5)</td>
<td>2 (9.5)</td>
<td>3 (4.7)</td>
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<tr>
<td>Fingolimod</td>
<td>11 (5.4)</td>
<td>1 (5.3)</td>
<td>1 (3.7)</td>
<td>1 (4)</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>1 (4.8)</td>
<td>7 (10.9)</td>
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<td>Glatiramer acetate</td>
<td>1 (0.5)</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>1 (1.6)</td>
</tr>
<tr>
<td>Natalizumab</td>
<td>6 (3.0)</td>
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<td>—</td>
<td>—</td>
<td>1 (5.9)</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>5 (7.8)</td>
</tr>
</tbody>
</table>

Abbreviations: DMT = disease-modifying therapy; EDSS = Expanded Disability Status Scale; IQR = interquartile range; NfL = neurofilament light chain; PMS = progressive MS; RRMS = relapsing-remitting MS.

a NfL only, MRI only, and clinical only correspond to patients with MS who had only elevated CSF NfL, clinical activity, or MRI activity, respectively. Combined disease activity groups are also provided.
b Each characteristic’s percentage represents their respective weight among the total patients in each disease activity group.
baseline EDSS score of 6 (IQR 4–6.5). Distribution of demographic characteristics by disease activity status is summarized in table 1.

In patients with PMS, 62 of 86 (72%) had treatment strategies including elevated CSF NfL measurements in the process, and in 17 (20%) an elevated CSF NfL measurement was the only evidence of disease activity used in the treatment decision-making process. In patients with RRMS, 75 of 117 (64%) had treatment strategies integrating elevated CSF NfL measurements with other measures, and in 5 (4%), elevated CSF NfL measurements were the only disease activity criterion used in the treatment decision-making process.

Treatment strategies consisted of No Treatment/No Escalation decisions in 62 patients (30.5%) and Treatment/Escalation decisions in 141 patients (69.5%) (table 2). Higher median CSF NfL measurements were observed for patients ultimately receiving a Treatment/Escalation decision (612 pg/mL) compared with No Treatment/No Escalation patients (264.5 pg/mL) (p < 0.001) (table 2, figure 3A). In the No Treatment/No Escalation group, there was little difference between median CSF NfL measurements of patients for whom we decided not to treat (264.5 pg/mL) compared with those whose treatment was not escalated (269 pg/mL). In the Treatment/Escalation group, there was a stepwise increase in median CSF NfL values from patients put on first-line injectable and oral DMTs (369 pg/mL), to those started on highly active DMTs (411 pg/mL), and those whose treatment was escalated (696 pg/mL).

The impact of CSF NfL-based treatment strategies on disability outcomes

Of the 95 patients in the No Treatment/No escalation group, the median baseline EDSS score was 2.5 (IQR 1.5–6), and the median follow-up EDSS score was 2.5 (IQR 1.5–6). Of the 108 patients in the Treatment/Escalation group, the median baseline EDSS score was 4 (IQR 1.75–6), and the median follow-up EDSS score was 4.75 (IQR 2–6). The median EDSS change was 0 (IQR 0–0.5) in both the No Treatment/No Escalation and the Treatment/Escalation group. The change in the EDSS was not found to be different at follow-up between the 2 treatment groups (p = 0.81) (figure 3B).

CSF NfL test experience (December 2015–July 2018)

The learning effect and adoption of the CSF NfL test was evaluated over the period of December 2015 through to July 2018 based on clinical documentation. The proportion of Treatment/Escalation vs No Treatment/No Escalation strategies did not differ from year to year in patients with elevated CSF NfL measurements (p = 0.99) (figure S1A, links.lww.com/NXI/A307); while in patients with normal CSF NfL measurements the proportion of No Treatment/No Escalation strategies significantly increased toward 2018 (p < 0.02) (figure S1B).

Discussion

In the present study, we evaluated the use of CSF NfL measurements to assist treatment strategies for patients with MS in clinical practice. Although CSF NfL has gained traction in research as a tool to identify disease activity in MS, it is not known whether the same applies for clinical practice, particularly that of personalized care. We explored the utility of the CSF NfL test in real-life practice.

In our cohort, we found that CSF NfL measurements were elevated in 42% of samples tested. Although CSF NfL closely aligned with clinical disease activity and MRI activity, all 3 parameters were present together in only a small proportion of patients (15%). CSF NfL measurements have been shown to reflect both inflammatory and neurodegenerative components of MS,8,9,12,20,21 which is supported by our finding that CSF NfL was not only associated with relapses and Gd-enhancing lesions on MRI but also with our composites of clinical and radiologic disease activity. However, modeling CSF NfL with the other 2 disease activity parameters showed that these associations were largely independent of each other. Recently, both CSF and serum NfL measurements have been shown to correlate not only with current but also with previous clinical and MRI activity in RRMS.22,23 Although assessments of disease activity (i.e., CSF NfL testing and clinical and radiologic assessments) were deemed to take place within a similar time frame, no predefined time intervals were set for this real-word study. This may explain the lack of association between CSF NfL measurements and some clinical and MRI measures evaluated in our study. The reasons for the association between sex and CSF NfL measurements found in our study remain uncertain. However, our findings are in line with the results of other studies showing that CSF NfL levels are significantly higher in men compared with...
women in healthy individuals and several neurodegenerative diseases.24

Most of the patients demonstrating an elevated CSF NfL measurement as the only evidence of disease activity had a progressive disease course. This observation was less apparent when looking at the clinical activity only or MRI activity only subgroups. Similarly, an elevated CSF NfL measurement was the only evidence of disease activity used in the treatment decision-making process in one-fifth of the patients with PMS within this cohort. This suggests that there is a subgroup of patients with MS with disease activity that may be missed by relying on clinical activity or MRI disease activity markers alone.7,25 Although a number of publications report on elevated CSF NfL during relapse in RRMS, subclinical inflammatory disease activity in PMS is often overlooked.7 Elevated CSF NfL in isolation may therefore still be acted on and has the potential to be much more than an aid for prognostication or treatment response.

Although CSF NfL measurements were only 1 of the 3 pillars in assessing disease activity during the treatment decision-making process, it is associated with clinical and radiologic activity as shown in Figure 2. The box plots demonstrate the relationship between CSF NfL (n = 203) and (A) MRI activity (new/enlarging lesions and/or Gd-enhancing lesions), (B) Gd-enhancing lesions, (C) clinical activity (relapses and/or progression), and (D) relapses. (E) Regression modeling NfL measurements as a function of clinical activity (relapses and/or progression) and MRI activity (new/enlarging lesions and/or Gd-enhancing lesions). Box-whisker plots represent median, quartiles, and 1.5 × interquartile range. Gd = gadolinium; NfL = neurofilament light chain.
making process, patients with elevated CSF NfL were more likely to be directed toward Treatment/Escalation strategies in our study. The EDSS outcomes at follow-up between the No Treatment/No Escalation and Treatment/Escalation strategies did not differ significantly in this study, suggesting counterpoise. This is in line with current evidence that more aggressive treatment strategies lead to better clinical outcomes and higher rates of NEDA.5,26 Moreover, early treatment in MS has been found to reduce the risk of converting from RRMS to PMS.27–29 As CSF measurements of NfL predict future disability progression in patients with MS,11 adding CSF NfL measurements in the workup of patients with MS, starting with the diagnostic lumbar puncture, may help refine treatment strategies (highly active vs first-line therapies).

A key hurdle for any new biomarker or treatment is the adoption in clinical practice, which is dependent on evidence combined with the willingness and feasibility to introduce the novelty.30 In our center, we did not detect a meaningful change in treatment strategies in those with elevated CSF NfL over time, whereas more No Treatment/No Escalation decisions were documented in those with normal CSF NfL measurements toward the end of the study period. Arguably, although an important reason for the adoption of CSF NfL has been to identify patients with

### Table 2 Treatment strategies according to CSF NfL measurements

<table>
<thead>
<tr>
<th>Treatment strategy</th>
<th>Elevated CSF NfL measurements (n = 85)</th>
<th>Normal CSF NfL measurements (n = 118)</th>
<th>All patients (N = 203)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>No. (%)</td>
<td>Median CSF NfL (pg/mL)</td>
<td>No. (%)</td>
</tr>
<tr>
<td>No treatment/escalation</td>
<td>4 (4.7)</td>
<td>701</td>
<td>58 (49.2)</td>
</tr>
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<td>No treatment</td>
<td>1 (1.2)</td>
<td>861</td>
<td>35 (29.7)</td>
</tr>
<tr>
<td>No escalation of treatment</td>
<td>3 (3.5)</td>
<td>453</td>
<td>23 (19.4)</td>
</tr>
<tr>
<td>Treatment/escalation, no. (%)</td>
<td>81 (95.3)</td>
<td>971</td>
<td>60 (50.8)</td>
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<tr>
<td>First-line injectable and oral DMTs</td>
<td>15 (17.6)</td>
<td>808</td>
<td>18 (15.3)</td>
</tr>
<tr>
<td>Highly active DMTs</td>
<td>61 (71.8)</td>
<td>996</td>
<td>35 (29.7)</td>
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<tr>
<td>Treatment escalation</td>
<td>5 (5.9)</td>
<td>682</td>
<td>7 (5.9)</td>
</tr>
</tbody>
</table>

Abbreviations: DMT = disease-modifying therapy; NfL = neurofilament light chain.

### Figure 3 The influence of CSF NfL on treatment strategies and disability outcomes

Box plots representing (A) the influence of CSF NfL on No Treatment/No Escalation vs Treatment/Escalation strategies. The horizontal black line indicates the age-specific reference value for CSF NfL (<30: 290 pg/mL, 30–39: 380 pg/mL, 40–59: 830 pg/mL). (B) EDSS change at 1-year follow-up as a function of treatment strategies (i.e., No Treatment/No Escalation vs Treatment/Escalation) (n = 203). Box-whisker plots represent median, quartiles, and 1.5 × interquartile range. p Values were obtained using the Kruskal-Wallis test. EDSS = Expanded Disability Status Scale; NfL = neurofilament light chain.
active disease for DMT escalation, the test might also have the beneficial effect of reducing the escalation to highly active treatments in those with lower disease activity. However, more detailed qualitative work is needed to explore this in greater detail.

The main limitation of our study is the observational nature of the recorded data. Data regarding treatment strategies were retrieved from the electronic medical records and are therefore prone to reporting bias. Therefore, the influence of comorbidities, intolerance, or poor adherence to prior treatments and preferences of patients and neurologists on treatment strategies could not be accounted for in our analysis. An additional limitation of our study may lie in our categorization of treatment strategies, which was based on NHS England treatment algorithms and local policies. Some of the patients with higher levels of disability received off-label subcutaneous cladribine. Therefore, our results may not always be generalizable to practices elsewhere. We further acknowledge that normative data for CSF NfL age-related cutoffs were based on a small group of individuals. However, CSF collection is relatively invasive precluding the conduction of large-scale validation studies with healthy volunteers. Moreover, CSF NfL age-related cutoffs reported by Uman-Diagnostics are in line with CSF NfL measurements reported by others and have also been used in different research settings. Finally, this was not a randomized study, and we therefore acknowledge that the differences of baseline characteristics between groups may not be randomly distributed. As such, future studies should aim to balance groups to prevent such differences.

Taken together, the findings from this study demonstrate that CSF NfL measurements can be adopted in routine clinical practice in MS. They complement established markers of disease activity to guide treatment strategies. The test may have a specific utility in patients with PMS where both clinical and MRI activity are more likely to be stable. The CSF NfL test has been adopted into our practice and has an impact on clinical outcomes based on EDSS progression.

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Publication history

Appendix Authors

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<th>Location</th>
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References


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