Neurofilament Light Chain Levels Are Predictive of Clinical Conversion in Radiologically Isolated Syndrome

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Abstract

Background and Objectives
To evaluate the predictive value of serum neurofilament light chain (sNfL) and CSF NfL (cNfL) in patients with radiologically isolated syndrome (RIS) for evidence of disease activity (EDA) and clinical conversion (CC).

Methods
sNfL and cNfL were measured at RIS diagnosis by single-molecule array (Simoa). The risk of EDA and CC according to sNfL and cNfL was evaluated using the Kaplan-Meier analysis and multivariate Cox regression models including age, spinal cord (SC) or infratentorial lesions, oligoclonal bands, CSF chitinase 3–like protein 1, and CSF white blood cells.

Results
Sixty-one patients with RIS were included. At diagnosis, sNfL and cNfL were correlated (Spearman r = 0.78, p < 0.001). During follow-up, 47 patients with RIS showed EDA and 36 patients showed CC (median time 12.6 months, 1–86). When compared with low levels, medium and high cNfL (>260 pg/mL) and sNfL (>5.0 pg/mL) levels were predictive of EDA (log rank, p < 0.01 and p = 0.02, respectively). Medium-high cNfL levels were predictive of CC (log rank, p < 0.01). In Cox regression models, cNfL and sNfL were independent factors of EDA, while SC lesions, cNfL, and sNfL were independent factors of CC.

Discussion
cNfL >260 pg/mL and sNfL >5.0 pg/mL at diagnosis are independent predictive factors of EDA and CC in RIS. Although cNfL predicts disease activity better, sNfL is more accessible than cNfL and can be considered when a lumbar puncture is not performed.

Classification of Evidence
This study provides Class II evidence that in people with radiologic isolated syndrome (RIS), initial serum and CSF NfL levels are associated with subsequent evidence of disease activity or clinical conversion.

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Radiologically isolated syndrome (RIS) is a preclinical stage of multiple sclerosis (MS) characterized by MRI brain lesions typical of MS detected in patients with clinical conditions not suggestive of MS.1 Given the absence of clinical features suggestive of MS, the gold standard for defining RIS is the specific dissemination in space (DIS) criteria (3 or 4 2005 DIS criteria for diagnosis of MS), which have not been updated since 2009 despite changes in MS criteria.1-3 Natural history studies suggest that the clinical course for patients with RIS is heterogeneous.4,5 Not all individuals will experience definable clinical symptoms attributable to inflammatory CNS demyelination and axonal damage. Identifying patients at high risk of clinical conversion and disability over time is mandatory for optimal clinical care. The risk for a future event may be stratified by key risk factors, but no biological biomarkers are used in clinical practice.4-6 Retrospective studies performed by an international working group, the Radiologically Isolated Syndrome Consortium (RISC), have demonstrated that one-third of patients with RIS converted to MS after 5 years and that another one-third experienced new brain lesions on follow-up scans.4 After 10 years, most patients with RIS convert to MS.5 Several factors have been associated with the risk of clinical conversion (CC) to MS in RIS: male sex, younger age, positive CSF for oligoclonal bands (OCBs), infratentorial (IT) lesions, spinal cord (SC) lesions on the index scan, and evidence of disease activity (EDA) defined by gadolinium-enhancing (Gd+) lesions on MRI follow-up.1,4,5,7,8 These prognostic factors have been confirmed on a large prospective cohort who had Gd+ lesions on the index scan.6

Several additional biomarkers have been explored for their specificity for MS. These include MRI markers such as the central vein sign that reflects perivenous inflammatory demyelination and can help differentiate MS from other white matter disorders.9 Another is the paramagnetic rim sign, which can be detected in most patients with RIS, suggesting the presence of subclinical chronically active demyelination at an early stage of the disease.10 Visual evoked potential anomalies and peripapillary retinal nerve fiber layer thickness measured by optical coherence tomography have also been associated with the risk of conversion to MS.7,11 Biological markers of MS in patients with RIS include CSF OCBs, high CSF neurofilament light chain (cNfL) levels, and IL-8 levels. By contrast, high CSF chitinase 3–like protein 1 (CHI3L1) and high CSF IL-17 levels are not associated with the risk of a clinical event.12-15 However, no peripheral biological marker providing additional prognostic value to epidemiologic, MRI, and CSF markers has so far been identified.16 CSF is a fluid of choice for characterizing biomarkers of brain disorders and has been widely investigated in MS at all stages of the disease.17,18 cNfL is a useful biomarker in several inflammatory and neurodegenerative CNS diseases.19 In MS, cNfL was mainly associated with EDA in relapsing-remitting MS (RRMS), clinically isolated syndromes (CIS), and RIS.12,20,21

In this study, we explored the value of sNfL as a predictive biomarker of disease activity in patients with RIS. We evaluated the predictive value of sNfL and cNfL levels for EDA and CC in patients with RIS in the context of other prognostic factors.

**Methods**

This study followed the STARD 2015 (Standards for Reporting Diagnostic accuracy studies) reporting guideline.

**Standard Protocol Approvals, Registrations, and Patient Consents**

This study used biological samples previously withdrawn in a prospective study, which was approved by a local ethics committee with patient informed consent. Because this study retrospectively analyzed data from a cohort of patients, it was approved by the Institutional Review Board of Nîmes (IRB no. 22.01.09), and no participant consent was required, according to the French Law.22

**Patients**

We selected a multicenter retrospective cohort of patients with untreated RIS from University Hospital Centers of Nice, Istanbul, and Nîmes on the basis of availability of CSF and/or serum samples. CSF samples from 50 patients and serum samples from 57 patients had been prospectively collected (biobank registered under no. 914066V2) between 2001 and 2018 after discovering unexpected demyelinating lesions fulfilling Okuda criteria,1 processed locally and centralized for this study. Epidemiologic, MRI, and biological data had been collected from patients with RIS, and CC occurrence prospectively followed. Patients treated with disease-modifying
were performed at 4°C instead of room temperature.27 Aliquots
conducted, except that sample transport and centrifugation
at the end of lumbar punctures and centrifuged within 2 hours at 1,500
g. CSF samples were collected using polypropylene tubes at the
collection with immunoblotting.
determined during biobanking as a routine test in each center.
There was no standardized MRI protocol, but all patients had
1.5T or 3.0T brain and spinal cord MRI studies. For all brain
MRI studies, 3D FLAIR and 3D MPRAGE sequences were
acquired. Standard CSF analyses from local laboratories were
collected (red cell count, white blood cell [WBC] count, and
CSF protein concentration). The presence of OCBs was also
determined during biobanking as a routine test in each center
by parallel isoelectric focusing of serum and CSF in combina-
tion with immunoblotting.

CSF and Serum Sample Collection
CSF samples were collected using polypropylene tubes at the
end of lumbar punctures and centrifuged within 2 hours at 1,500
X g for 10 minutes according to the guidelines of the BioMS-eu
consortium, except that sample transport and centrifugation
were performed at 4°C instead of room temperature.27 Aliquots
(500 μL) were stored at −80°C in 1.5 mL tubes (Protein LoBind
0030108.116, Eppendorf) until use. Patients with traumatic
lumbar punctures (>500 red cells/mm³) were excluded.

Serum samples were spun at the latest 2 hours after blood
sampling at 1,500 X g for 10 minutes at room temperature and
aliquots (500 μL) stored at −80°C in 1.5 mL tubes (Protein LoBind
0030108.116, Eppendorf) until use.

Measure of CSF and Serum Biomarkers
All serum and CSF samples were centralized at the Clinical
Proteomics Platform of the Laboratory of Biochemistry—
Clinical Proteomic in Montpellier. sNfL and cNFL concentra-
tions were determined using commercial NF-Light assay
(Quanterix, Billerica, MA) based on ultrasensitive Simoa
technology, as previously described.28 CSF CHI3L1 levels
were measured by ELISA using the MicroVue YKL-40 EIA kit
(Quidel Corporation, San Diego, CA), as already described.29
All experiments were performed with a single batch of re-
agents by researchers blinded from any clinical information.
There were no missing values for NfL or CHI3L1 measures.

Statistical Analysis
Statistical analyses were performed using the R software (v3.0.2)
and SAS software, version 9.4 (SAS Institute, Cary, NC). The
type I error rate was 0.05, and no correction for multiple testing
was performed. Median NfL values were compared using the
nonparametric Mann-Whitney test. Area under the curve (AUC)
values were estimated from time-dependent ROC
curves for censored data using the inverse probability of cen-
soring weighting technique of Uno’s method30 for the occurrence
of events at 2, 3, and 5 years. The time from RIS diagnosis to CC
or EDA was analyzed using the Kaplan-Meier estimator and
compared between groups using the log-rank test. Because log
hazard of CC or EDA and the continuous value of cNFL and
sNfL did not have a linear relationship, we chose to perform an
analysis based on the tertiles of the distribution of these variables.
These analyses did not show a risk gradient between the me-
dium and high tertiles of cNFL and sNfL (see Results section
Figure 2), indicating a possible threshold effect. We therefore
continued our analyses by dichotomizing these variables on the
value of the first tertile and thus grouping the medium and high
tertiles. Age, sNfL (low vs medium-high), cNFL (low vs
medium-high), CSF CHI3L1, CSF WBCs, OCBs, 3 or 4 2005
DIS criteria, and the presence Gd+ lesions, IT lesions, and SC
lesions were tested in a univariate Cox model analysis, followed
by a multivariate analysis when significant at a threshold of 0.2 in
univariate analyses. Factors with more than 10% of missing data
were not included in the multivariate models. Finally, we an-
alyzed the additive value of sNfL and cNFL by comparing the
predictive value of several predictive models based on the SC
lesions and OCBs, with and without the addition of sNfL (low vs
medium-high) or cNFL (low vs medium-high), analyzing the
differences in Uno concordance statistic (for censored data)
between the different models.31

Classification of Evidence
This study is based on an RIS cohort study with prospective
data collection in local databases using EDMUS. RIS status
was determined centrally according to the presence of 2009
RIS criteria without neurologic symptoms of MS, without
knowledge of the diagnostic test results for OCBs, CSF
CHI3L1 levels, CSF WBC count, sNfL levels, and cNFL levels.
These markers were determined in more than 80% of the
participants. Patients were prospectively followed up from
RIS discovery with regular MRI and clinical evaluation.

Data Availability
Anonymized participant data that underlie the results reported
in this article will be shared beginning 6 months and
ending 12 months after article publication on reasonable re-
quest by qualified investigators.

Results
Patient Characteristics at Diagnosis
Sixty-one patients with RIS (75% females, mean age 37 years;
SD 12) with a biocollection of serum (n = 57) and/or CSF
(n = 50) samples were included in the study (Table 1). Of the
processed samples, 47% had CSF OCBs, median WBC count
was 2/mm³ (0–25), 23% had 4 2005 DIS criteria, 20% had at
least 1 brain Gd+ lesion, and 29% had at least 1 infratentorial
(II) lesion. In a subgroup of 59 patients with spinal cord
MRI, 47% had at least 1 SC lesion, none of which was en-
hanced after administration of gadolinium. Median (95% CI)
cNfL and sNfL levels were 383 (276; 569) pg/mL and 6.9 (5.3; 7.6) pg/mL, respectively, and median (95% CI) CSF CHI3L1 level was 135 (108; 183) ng/mL (Table 1).

In a subgroup of 46 patients (75%) with both sNfL and cNfL levels available, sNfL and cNfL levels were correlated (Spearman correlation coefficient r = 0.780, p < 0.01, Figure 1A). Neither sNfL (r = 0.098, p = 0.47) (Figure 1B) nor cNfL levels (r = 0.020, p = 0.89) were correlated with age (data not shown). sNfL levels were not significantly different between male and female patients, RIS patients with 3 or 4 2005 DIS criteria, and those with or without SC lesion (Figure 1, C and D). cNfL levels, but not sNfL levels, were significantly higher in RIS patients with positive OCBs than in those with negative OCBs (p = 0.02, Mann Whitney test, Figure 1, C and D).

Evolution of Patients With RIS During Follow-up

Patients were prospectively followed up from diagnosis of RIS for a median time of 23 months (2–228 months). During follow-up, 36 patients with RIS (59%) evolved to a CC with a median conversion time of 12.6 months (1–86 months), while 25 patients remained NC after a median follow-up of 78 months (7–228 months). Forty-seven patients with RIS (76%) showed EDA in a median delay of 11.1 months (1–86 months). The characteristics of the subgroups of patients regarding EDA vs NEDA and CC vs NC are summarized in Table 1. cNfL levels were higher in RIS patients with CC vs NC and EDA vs NEDA (464 pg/mL vs 253 pg/mL, p = 0.5, and 464 pg/mL vs 161 pg/mL, p < 0.01, respectively). sNfL levels were also higher in RIS patients with EDA vs NEDA (7.1 pg/mL vs and 4.0 pg/mL, p < 0.01), but not in RIS patients with CC compared with NC (7.2 vs 5.7 pg/mL, p = 0.13).

Association of cNfL and sNfL With Clinical Conversion and Evidence of Disease Activity

Using ROC curves, we determined time-dependent AUC [95% CI] values of sNfL and cNfL that best discriminate EDA vs NEDA and CC vs NC at 2, 3, and 5 years of follow-up (Table 2). Best AUC values for CC vs NC were provided at 3 years (0.76 [0.59–0.92] for cNfL and 0.66 [0.47–0.86] for sNfL). Best AUC values for EDA vs NEDA were provided at 5 years (0.89 [0.76–1.00] for cNfL and 0.78 [0.54–1.00] for sNfL).

Comparison of cNfL and sNfL AUC [95% CI] values for CC and EDA at 3 years in a subgroup of 46 patients with both cNfL and sNfL levels available showed that cNfL has a more robust association profile with CC (0.77 [0.61; 0.92] and 0.62 [0.41; 0.84], respectively) and EDA (0.82 [0.65; 0.98] and 0.69 [0.49; 0.88], respectively) than sNfL, although not significantly.

Individual Factors Predicting Clinical Conversion and Evidence of Disease Activity

Using cutoff values defined as the first tertile of distribution as cutoff values (cNfL >260 and sNfL >5.0 pg/mL) and taking into account the time of follow-up, the Kaplan-Meier analysis revealed that the presence of IT lesion(s) and OCBs, medium-high cNfL, and medium-high sNfL are predictive of EDA (log rank, p = 0.03, p = 0.03, p < 0.01, and p = 0.02,

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**Table 1 Characteristics of the Total RIS Population and Subgroups According to Evidence of Disease Activity or Clinical Conversion During Follow-up**

| Patient characteristics | RIS n = 61 | RIS-NC n = 25 | RIS-CC n = 36 | RIS-NEDA n = 14 | RIS-EDA n = 47 |
|-------------------------|------------|---------------|---------------|----------------|---------------|
| Sex ratio (female %)    | 75         | 76            | 75            | 79             | 74            |
| Mean age (y)            | 37         | 37            | 37            | 38             | 36            |
| ≥2 OCBs (%)             | 47         | 36            | 56            | 21             | 55            |
| Median CSF WBC count (/mm³) | 2         | 1             | 2             | 1              | 2             |
| Mean CSF WBC count (/mm³) | 3         | 2             | 4             | 2              | 4             |
| Four 2005 DIS criteria, % | 23     | 12            | 31            | 21             | 28            |
| Presence of IT lesion(s), % | 29     | 16            | 36            | 0              | 38            |
| Presence of brain Gd+ lesion(s), % | 20 | 28            | 14            | 21             | 19            |
| Presence of SC lesion(s) (n = 59), % | 47 | 17            | 67            | 21             | 53            |
| Presence of SC Gd+ lesion(s) (n = 59), % | 0 | 0             | 0             | 0              | 0             |
| Median CSF CHI3L1 levels (ng/mL) (n = 50) | 135 | 133           | 162           | 111            | 162           |
| Median cNfL levels (pg/mL) (n = 50) | 383 | 253           | 464           | 161            | 464           |
| Median sNfL levels (pg/mL) (n = 57) | 6.9  | 5.7           | 7.2           | 4.0            | 7.1           |

Abbreviations: CC = clinical conversion; CHI3L1 = chitinase 3–like protein 1; cNfL = CSF neurofilament light chain; DIS = dissemination in space; EDA = evidence of disease activity; Gd+ = gadolinium-enhancing; IT = infratentorial; NC = no clinical conversion; NEDA = nonevidence of disease activity; OCBs = oligoclonal bands; RIS = radiologically isolated syndrome; SC = spinal cord; sNfL = serum neurofilament light chain; WBCs = white blood cells.
respectively, Figure 2, A and C, Figure 3, A and C). It also indicated that the presence of SC lesion(s), medium-high cNfL, and elevated WBCs ($\geq 5$/mm$^3$) are predictive of CC (log rank, $p < 0.01$, $p < 0.01$, and $p = 0.04$, respectively, Figure 2, B and D, Figure 3B), while medium-high sNfL was not (log rank, $p = 0.08$, Figure 3D). Age, CSF CHI3L1, number of 2005 DIS criteria, and the presence of Gd+ lesions on the index MRI were not significantly associated with EDA nor CC (data not shown).

Cox univariate analysis of age, sNfL, cNfL, CSF CHI3L1, CSF WBCs, OCBs, number of 2005 DIS criteria, Gd+ lesions, IT lesions, and SC lesions on the index MRI revealed that medium-high cNfL and the presence of SC lesions were predictive of CC, while medium-high cNfL, medium-high sNfL, the presence of OCBs, and IT lesions were predictive of EDA (Table 3).

**Multivariate Analysis of Prognostic Factors in RIS**

cNfL and sNfL being highly correlated, we built 2 multivariate Cox regression models to identify EDA and CC prognostic factors in patients with RIS (Table 3).

In the multivariate model analyzing cNfL, medium-high cNfL was an independent predictive factor of EDA (HR = 8.0, $p <$
0.01). By contrast, medium-high cNfL and the presence of SC lesion(s) were independent factors of CC (HR = 4.8, \( p < 0.01 \) and HR = 2.7, \( p = 0.03 \), respectively, Table 3).

In the multivariate model analyzing sNfL, medium-high sNfL and SC lesions were independent predictive factors of EDA (HR = 3.0, \( p = 0.01 \) and HR = 2.2, \( p = 0.03 \), respectively),

### Table 2  Sensitivity, Specificity, and Area Under the Curve of sNfL and cNfL (Using First Tertile as Threshold for Positivity) for Evidence of Clinical Activity and Clinical Conversion at 2, 3, and 5 Years, Determined by Time-Dependent ROC Curves

|         | 2 y            | 3 y            | 5 y            |
|---------|----------------|----------------|----------------|
|         | Se (%)         | Sp (%)         | AUC (95% CI)   | Se (%)         | Sp (%)         | AUC (95% CI)   | Se (%)         | Sp (%)         | AUC (95% CI)   |
| cNfL    |                |                |                |                |                |                |                |                |                |
| CC      | 83.3           | 62.5           | 0.71 (0.53; 0.88) | 88.4           | 66.7           | 0.76 (0.59; 0.92) | 85.5           | 58.3           | 0.68 (0.45; 0.92) |
| EDA     | 82.1           | 78.9           | 0.79 (0.62; 0.97) | 87.1           | 75.0           | 0.81 (0.66; 0.96) | 85.9           | 87.5           | 0.89 (0.76; 1.00) |
| sNfL    |                |                |                |                |                |                |                |                |                |
| CC      | 73.1           | 60.7           | 0.64 (0.47; 0.82) | 67.7           | 68.0           | 0.66 (0.47; 0.86) | 87.2           | 50.0           | 0.62 (0.42; 0.82) |
| EDA     | 67.7           | 69.6           | 0.67 (0.49; 0.85) | 73.5           | 75.0           | 0.72 (0.55; 0.90) | 87.3           | 66.7           | 0.78 (0.54; 1.00) |

Abbreviations: AUC = area under the curve; CC = clinical conversion; cNfL = CSF neurofilament light chain; EDA = evidence of disease activity; Se = sensitivity; sNfL = serum neurofilament light chain; Sp = specificity.
while medium-high sNfL, the presence of SC lesion(s), and OCBs were independent factors of CC (HR = 2.8, $p = 0.02$, HR = 3.3, $p < 0.01$, and HR = 2.36, $p = 0.02$, respectively) (Table 3). Especially, 84% of patients with RIS with medium-high sNfL levels, SC lesions, and OCBs had a CC at 3 years, compared with 34% in other patients with RIS.

To analyze the additive value of NfL to SC lesions and OCBs, we further compared the AUC values of SC lesions + OCBs and SC lesions + OCBs + NfL for EDA and CC at 3 years (Table 4). All AUCs values were improved when adding sNfL or cNfL to SC lesions and OCBs for the prediction of EDA and CC, although it was statistically significant only for cNfL and EDA at 3 years (AUC = 0.65 [0.43; 0.86] for SC lesions + OCBs, vs AUC = 0.85 [0.67; 1.00] for SC lesions + OCBs + cNfL, $p = 0.01$ for AUC comparison).

**Classification of Evidence**

This study provides Class II evidence that in people with RIS, initial serum and CSF NfL levels are associated with subsequent evidence of disease activity or clinical conversion.

**Discussion**

We compared the value of NfL levels in serum and CSF samples from patients with RIS in the context of clinical risk. We demonstrated that sNfL and cNfL levels were highly correlated and constituted independent predictors of CC. Our results align with the most extensive RIS studies that described the importance of SC lesions for RIS prognosis, also identified here as an independent predictor of CC in the multivariate Cox model. These factors reflect the risk of CC during follow-up. They could be considered to counsel patients with RIS during
diagnosis and to organize the clinical and MRI surveillance to detect disease activity. Indeed, EDA is a classical endpoint in the follow-up of patients with MS, but predictive factors of EDA have never been investigated in patients with RIS. Identifying OCBs and SC lesions as predictors of EDA also suggests that these factors provide complementary information to sNfL and cNfL that remain independent predictive factors in both cases (EDA and CC). Moreover, sNfL or cNfL provide additive value to SC lesions and OCBs to discriminate RIS patients with CC or EDA from clinically and/or radiologically stable RIS patients, especially for cNfL and EDA (Table 4). As a matter of fact, NfL are released from axons and reflect neuroaxonal injury in MS. NfLs seem to be especially associated with focal inflammation (active plaques and relapses) than to the neurodegenerative process associated with disease progression.32 Recently, improved detection of sNfL based on ultrasensitive ELISA kits or Simoa allowed to reliably measure sNfL concentrations that are well correlated to that of CSF in patients with MS.33 The high sensitivity of this test, illustrated by the

| Table 3: Analysis of Baseline Characteristics Predictive of Evidence of Disease Activity or Clinical Conversion During Follow-up Using Multivariate Cox Models |

| Variable | Univariate analysis | Multivariate analysis |
|----------|---------------------|----------------------|
|          | HR | 95% CI | p Value | HR | 95% CI | p Value |
| Time to CC | cNfL | 3.9 | 1.6; 9.8 | <0.01 | cNfL | 4.8 | 1.8; 12.8 | <0.01 |
|          | sNfL | 2.1 | 0.4; 4.5 | 0.07 | SC | 2.7 | 1.1; 6.7 | 0.03 |
|          | SC | 3.3 | 1.6; 6.6 | <0.01 | OCBs | 1.8 | 0.8; 3.8 | 0.13 |
|          | OCBs | 1.8 | 0.9; 3.5 | 0.08 | Gd+ | 0.6 | 0.2; 1.8 | 0.33 |
|          | Gd+ | 0.4 | 0.2; 1.2 | 0.10 |              | 0.7 | 0.3; 1.7 | 0.47 |
|          | 2005 DIS | 1.6 | 0.8; 3.3 | 0.18 | sNfL | 2.8 | 1.2; 6.6 | 0.02 |
|          | WBCs | 1.9 | 1.0; 3.9 | 0.07 | SC | 3.3 | 1.4; 7.7 | <0.01 |
|          | IT | 1.5 | 0.7; 2.9 | 0.28 | OCBs | 2.4 | 1.1; 4.9 | 0.02 |
|          | Age | 1.0 | 0.5; 1.9 | 0.89 | Gd+ | 0.4 | 0.1; 1.7 | 0.22 |
|          | CHI3L1 | 15 | 0.8; 3.0 | 0213 | 2005 DIS | 13 | 06; 2.7 | 055 |
| Time to EDA | cNfL | 5.9 | 2.5; 13.8 | <0.01 | cNfL | 8.0 | 2.8; 22.7 | <0.01 |
|          | sNfL | 2.2 | 1.1; 4.5 | 0.02 | OCBs | 1.3 | 0.6; 2.7 | 0.43 |
|          | OCBs | 2.1 | 1.2; 3.7 | 0.01 | SC | 1.8 | 0.9; 3.9 | 0.12 |
|          | SC | 1.6 | 0.9; 2.9 | 0.12 | CHI3L1 | 1.0 | 0.5; 2.1 | 0.99 |
|          | CHI3L1 | 1.6 | 0.9; 3.0 | 0.12 | IT | 1.0 | 0.5; 2.1 | 0.96 |
|          | IT | 2.1 | 1.2; 3.9 | 0.01 | sNfL | 3.0 | 1.3; 6.8 | 0.01 |
|          | 2005 DIS | 1.5 | 0.8; 2.9 | 0.19 | OCBs | 1.7 | 0.8; 3.8 | 0.17 |
|          | Gd+ | 0.8 | 0.4; 1.7 | 0.55 | SC | 2.2 | 1.1; 3.8 | 0.03 |
|          | Age | 0.9 | 0.5; 1.7 | 0.78 | CHI3L1 | 1.1 | 0.5; 2.4 | 0.75 |
|          | WBCs | 1.0 | 0.5; 2.0 | 0.94 | IT | 1.3 | 0.6; 3.0 | 0.45 |

Abbreviations: 2005 DIS = presence of four 2005 DIS criteria; CHI3L1 = chitinase 3-like protein 1; cNfL = CSF neurofilament light chain; EDA = evidence of disease activity; Gd+ = presence of brain gadolinium-enhanced lesion(s); HR = hazard ratio; IT = infratentorial; LCC = clinical conversion; NfL = neurofilament light chain; OCBs = oligoclonal bands; SC = presence of spinal cord lesion(s); sNfL = serum neurofilament light chain; sNfL = serum NfL; WBCs = white blood cells.

Items with a p value <0.2 were included in the multivariate Cox regression models and those with a p value <0.05 are in bold.

* cNfL >260 pg/mL.
* sNfL >5.0 pg/mL.
* WBCs >5/mm3.
* Age older than 37 years.
* CSF CHI3L1 > 135 ng/mL.
absence of missing values, allows accurate measures of sNfL in serum. This study provides the first measurement of sNfL in patients with RIS using the NF-light assay. Using this assay, the median sNFL (6.9 pg/mL) in patients with RIS was lower than that observed by others in patients with presymptomatic MS (16.7 pg/mL) and patients with CIS (17.0 pg/mL) using the same antibody adapted to Simoa homebrew kits. However, our results corroborate sNfL median levels measured in our previous study using the NF-light assay in patients with MS (9.4 pg/mL) and those recently found in patients with early RRMS (10.1 pg/mL) using the same assay.

Compared with the electrochemiluminescence-based method or ELISA, the high accuracy of the Simoa assay, even at lower values, allows sNfL to strongly correlate with cNfL. This probably contributed to the performance of sNfL as a prognostic biomarker of clinical evolution in patients with RIS in our study. However, cNfL showed an overall more robust association profile with CC and EDA, suggesting that cNfL better reflects disease activity and predicts the evolution of patients with RIS than sNfL. This might be the consequence of differences among individuals in the kinetics of neurofilament protein release from neurons and trafficking between the brain and blood compartments. Nevertheless, blood sampling is less invasive, making sNfL an excellent alternative to cNfL for RIS prognosis when a lumbar puncture cannot be performed. More extensive validation studies are required before sNfL may replace cNfL.

The high accuracy of the Simoa assay probably enabled sNfL to overcome OCBs to predict CC. Even if this suggests that OCBs might no longer be required for the prognosis of CC in RIS patients with SC lesions, their detection provides a reasonable specificity for the differential diagnosis of RIS vs other conditions. Contrasting with the observations of a previous study showing that OCBs and cNfL were independent predictive factors of CC in patients with RIS, OCBs and sNfL were independent predictors of CC in our study. Including the presence of SC lesions, one of the most robust predictors of CC in our Cox regression analysis might explain why cNfL and SC lesions, but not OCBs, were independent predictors of CC in our study. This discrepancy might also reflect the different populations, time of follow-up, and assays used to measure NfL (ELISA vs Simoa) or the association between the presence of OCBs and cNfL levels that potentially minimize their prognostic value (Figure 1D). Nevertheless, we confirm that CSF CHI3L1 is not a predictor of EDA and CC in patients with RIS.

This study has some limitations given its retrospective nature. Incomplete availability of CSF and serum samples to determine cNfL, CSF CHI3L1, and sNfL led to missing data, lowering the power of the regression models, limited to 50 and 57 patients for the cNfL and the sNfL models, respectively (Table 3). Moreover, the small size and the small number of centers limit knowledge regarding how these data are translatable to other races and ethnicities with RIS. In addition, although our population was comparable with cohorts from previous published RIS studies (age, sex ratio, MRI, and CSF findings), the proportion of patients with RIS converting to MS was higher (50% converters at 36 months) in our cohort. This might be explained by the selection of patients with untreated RIS only. This also probably enhanced the statistical power of the study for identifying predictors of CC and constitutes a potential caveat for future confirmation studies with less active cohorts. As illustrated in the Kaplan-Meier analyses, cNfL and sNfL values did not

### Table 4 Additive Value of sNfL and cNfL to SC Lesions and OCBs to Discriminate CC and EDA From Clinically and/or Radiologically Stable RIS Patients at 3 Years

| Endpoint | Combinations of variables | Se (%) | Sp (%) | AUC (95% CI) | AUC comparison p Value |
|----------|---------------------------|--------|--------|--------------|------------------------|
| **cNfL (n = 50)** | SC lesion + OCBs | 70.8 | 68.0 | 0.77 (0.66; 0.88) | 0.46 |
| | SC lesion + OCBs + cNFL | 66.7 | 78.5 | 0.79 (0.64; 0.94) | 0.65 |
| **EDA** | SC lesion + OCBs | 47.3 | 84.8 | 0.71 (0.56; 0.86) | 0.01 |
| | SC lesion + OCBs + sNFL | 78.9 | 69.7 | 0.77 (0.61; 0.93) | 0.01 |
| **sNfL (n = 57)** | SC lesion + OCBs | 40.0 | 89.0 | 0.70 (0.55; 0.86) | 0.13 |
| | SC lesion + OCBs + cNFL | 80.0 | 81.0 | 0.85 (0.71; 0.99) | 0.01 |
| **EDA** | SC lesion + OCBs | 66.7 | 61.0 | 0.65 (0.43; 0.86) | 0.01 |
| | SC lesion + OCBs + sNFL | 93.3 | 77.4 | 0.85 (0.67; 1.00) | 0.01 |

Items with a p value <0.05 are in bold.

AUC = area under the curve (Uno Concordance Statistic); CC = clinical conversion; cNFL = CSF neurofilament light chain >260 pg/mL; EDA = evidence of disease activity; OCBs = positive oligoclonal bands; RIS = radiologically isolated syndrome; SC lesion = presence of spinal cord lesion(s); se = sensitivity; sNFL = serum neurofilament light chain >5.0 pg/mL; sp = specificity.

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appear as proportionally correlated with CC and EDA risk in a linear relationship, with similar risks of conversion for medium values and high values of each marker, when compared with low values (Figure 2). This suggests that NfL increase above a threshold value of 260 pg/mL for cNfL and 5.0 pg/mL for sNfL (corresponding to the first tertile of cNfL and sNfL in our study) indicates active neuroaxonal injury predictive of EDA and CC, especially for cNfL (Table 3). Intriguingly, the cutoff value for cNfL to predict CC was lower using Simoa (260 pg/mL) than that previously determined by ELISA in patients with RIS (619 pg/mL). It can be the consequence of different assays used to measure NfL (ELISA vs Simoa) and of diverse populations investigated, as already observed in previous studies dedicated to MS biomarkers, such as those examining CHI3L1 in patients with CIS. A threshold value determined using Simoa of 5.0 pg/mL could be chosen for prospective studies aiming at validating the performance of sNfL alone or in combination with SC lesions and OCBs to predict CC and EDA and at establishing an RIS conversion score combining these factors.

In conclusion, we showed that elevated cNfL and sNfL levels at diagnosis are both predictive factors of EDA and CC in RIS. Our study provides the first analysis of sNfL in an RIS cohort and offers Class I evidence that sNfL >5.0 pg/mL is an independent predictor of CC and EDA in patients with RIS. Although cNfL predicts disease activity better, sNfL is more accessible and can be considered to identify patients at higher risk of conversion to help clinicians for counseling and monitoring patients when a lumbar puncture is not performed.

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