Biomarkers For MS

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Disclosures

None
Learning Objectives

Process of biomarker development

Biomarkers in clinical practice

Serum NfL as potential biomarkers
Key Messages

- Serum NfL levels are elevated in MS patients
- Serum NfL is not specific to MS and it increases gradually with normal ageing
- Levels correlate with number of T2 and GD+ lesions on MRI
- Baseline sNfL can predict Disability progression and change in brain volume after 5 –ears or more
Characteristics of an Ideal Biologic Marker

- Measurable in a body fluid that is easy to obtain
- Easily measurable with routine /affordable laboratory procedures
- Involved in the disease pathogenesis
- Correlated with clinical disease activity and disability
- High sensitivity in detecting relevant disease activity
- High specificity (not influenced by other diseases, complications such as infection)
- Correlated with radiological disease activity markers, such as MRI
- Undergo rapid normalization under therapy in responders
- Undergo no normalization under therapy in non-responders
Schematic Representation Of The Process Of Biomarker Development
Types Of Biomarker Needed In MS

- Supporting diagnosis
- Identify converters from CIS to MS
- Identify converters from RR to SPMS
- Predicting disease severity
- Predicting response to treatment
- Predicting risk of adverse events
Figure 1: Types of biomarkers in MS
<table>
<thead>
<tr>
<th>Category</th>
<th>Body fluid</th>
<th>Biomarker test</th>
<th>Target population</th>
<th>Prevalence</th>
<th>Effect on clinical decision making?</th>
</tr>
</thead>
<tbody>
<tr>
<td>Anti-natalizumab antibodies</td>
<td>Serum</td>
<td>ELISA</td>
<td>Patients with MS given natalizumab</td>
<td>4–1–6% of natalizumab-treated patients†‡§</td>
<td>Yes; if persistently positive, treatment with natalizumab should be discontinued§</td>
</tr>
<tr>
<td>Neutralising antibodies</td>
<td>Serum*, PBMCs</td>
<td>CPE*, luciferase gene-reporter assay*, MxA-based assays</td>
<td>Patients with MS given interferon beta</td>
<td>2–45% of patients given interferon beta†</td>
<td>Yes; if present, a switch to a non-interferon-beta treatment should be considered‡‡§</td>
</tr>
<tr>
<td>IgG oligoclonal bands</td>
<td>Diagnostic</td>
<td>CSF, serum</td>
<td>Patients suspected to have demyelinating disease</td>
<td>&gt;95% of patients with MS§</td>
<td>No</td>
</tr>
<tr>
<td>IgG index</td>
<td>Diagnostic</td>
<td>CSF, serum</td>
<td>Patients suspected to have demyelinating disease</td>
<td>Increased in 70% of patients with MS§</td>
<td>No</td>
</tr>
<tr>
<td>Anti-aquaporin-4 antibodies</td>
<td>Diagnostic</td>
<td>Serum*, CSF</td>
<td>Patients with clinical and MRI features suggestive of neuromyelitis optica or neuromyelitis optica spectrum disorders</td>
<td>Almost 75–90% of patients with neuromyelitis optica; almost absent in MS patients§</td>
<td>Yes; if present, diagnosis should be separate from typical MS‡‡‡</td>
</tr>
<tr>
<td>Anti-JC virus antibodies</td>
<td>Diagnostic</td>
<td>Serum*, plasma</td>
<td>ELISA</td>
<td>50–60% of patients with MS‡‡‡</td>
<td>Yes; test result allows estimation of the patient’s risk for progressive multifocal leukoencephalopathy‡§</td>
</tr>
<tr>
<td>Anti-VZV antibodies</td>
<td>Diagnostic</td>
<td>Serum*, plasma</td>
<td>ELISA</td>
<td>90–95% of patients with MS‡‡‡</td>
<td>Yes; seronegative patients should be vaccinated at least 1 month before start of fingolimod‡§§</td>
</tr>
</tbody>
</table>

Potential Biomarkers That Have Shown Some Replication Across Studies

- Neurofilament light in CSF and serum
- MOG antibodies
- Chitinase 3-like 1 and Chitinase 3-like 2
- Certain miRNAs
- Low affinity serum antibody microarrays
CH3L1 is a member of the family of chitinases and chitinase-like proteins containing a highly conserved glyco-18 domain as common feature. For these proteins, chitin is the only documented substrate.

CHI3L1 can bind chitin but lacks chitinolytic activity.

In the CNS, CHI3L1 expression has been mainly observed in astrocytes of monkeys and humans with lentiviral encephalitis, and patients with brain infarcts.
CSF Chitinase 3-like 1 Levels Are Associated With Conversion To MS

Brain 2010: 133; 1082–1093
Neurofilaments (NF)
Neurofilaments (NF)

NfH = neurofilament heavy chain; NfL = neurofilament light chain; NfM = neurofilament intermediate chain; N = N-terminus; C = C-terminus.
- Neurofilaments could be used in clinical practice as surrogate endpoints of neuroaxonal damage

- CSF concentrations of both NF-L and NF-H have been noted to be increased in patients with MS:
  - during relapse
  - related to radiological activity
  - related to disability and conversion to SPMS in 14 year follow up studies
  - noted to have prognostic value for conversion to MS in CIS patients-1yr

- NF-L seems to be a more accurate sign of acute axonal damage associated with inflammation than does NF-H

- NF-H better captures chronic axonal damage and shows a stronger association with disability progression

- A reduction in CSF NF-L is associated with MS treatments such as natalizumab, mitoxantrone, and rituximab, and NF-L has been proposed as a surrogate endpoint for treatment efficacy
Measurement of sNfL

CSF NfL is easily measured by ELISA, but serum Nfl are 70-100 X lower than CSF.

How do we measure sNfL:
- Conventional ELISA
- Electrochemiluminescence-based method (ECL assay)
- Single-molecule array (Simoa)

Analytical sensitivity was 0.62 pg/mL for Simoa, 15.6 pg/mL for the ECL assay, and 78.0 pg/mL for the ELISA.
Correlation Between Serum And CSF Levels Of NfL

Novakova et al. Neurol 2017;89:2230–2237

Disanto et al. ANN NEUROL 2017;81:857–870
sNfL Specificity

• Serum NFL level is a highly predictive marker of long-term poor neurologic outcome at 24 hours after cardiac arrest. *JAMA Neurol.* 2018 Oct 29

• Serum NFL levels were higher in ALS in comparison to other neurologic diseases except for CJD. A cut-off level of 62 pg/mL discriminated between ALS and all other conditions with 85.5% sensitivity (95% CI 78% to 91.2%) and 81.8% specificity (95% CI 74.9% to 87.4%). *J Neurol Neurosurg Psychiatry.* 2018 Oct 11. pii: jnnp-2018-318704.

• Serum NfL holds promise as a biomarker for monitoring primary and secondary neuroaxonal injury after ischemic stroke and for predicting functional outcome. *Neurology.* 2018 Sep 14

• serum NfL correlates with functional impairment and brain atrophy in bvFTD at different disease stages. *Neurology.* 2018 Sep 12
Effect of Ageing On sNfL

2.2% increase in sNfL for each additional year ($\beta = 1.022$, 95% CI=1.018–1.026, $p < 0.001$)

no statistically significant difference between males and females

no association between sNfL and storage time

Disanto et al. ANN NEUROL 2017;81:857–870
Association of sNfL at baseline with percentage change in brain volume over 5 years

Barro et al. BRAIN 2018: 141; 2382–2391
Association Between sNfL levels And MRI

Disanto et al. ANN NEUROL 2017;81:857–870
Association Between EDSS And sNfL

1-point EDSS increase corresponds to an sNfL increase of approximately 14.1%

Disanto et al. ANN NEUROL 2017;81:857–870
Change in sNfL After Treatment
Effect of fingolimod on NfL levels in blood, (A) compared with placebo, FREEDOMS study; (B) compared with interferon-β-1a, TRANSFORMS study
Irrespective of treatment, patients with high blood NfL concentrations (>60 pg/mL) at baseline compared with those with low baseline NfL concentrations (<30 pg/mL) had 2.6 times more new or enlarging T2 lesions (difference: 164%), 2.5 times more MS relapses (difference: 153%), 2.9 times more brain volume loss (difference: 195%) (all p < 0.001), and had a 1.9 times higher risk of 3-month CDW (p = 0.0605)
Conclusions

(1) sNfL levels can be reliably and reproducibly measured in serum samples from MS patients
(2) in independent HC and patient cohorts, sNfL levels are positively associated with age but not gender
(3) sNfL levels closely reflect NfL concentration in the CSF of MS patients
(4) sNfL levels are increased in MS patients as compared to HC and positively associated with T2 and GE lesions in both brain and spinal cord
(5) sNfL levels are increased in patients with recent relapses or worsening of disability, are higher with increasing EDSS scores, and decrease with increasing duration of DMT
(6) sNfL levels are associated with an increased risk of future relapses and EDSS worsening.
Questions that need to be answered in MS

Is it possible to monitor individual treatment responses via sNfL?

Can we predict neuronal loss or even transition into long-term progressive disease course in the very early stage of disease and match immunomodulatory therapy regimes?
What do we need before implementing sNfL measurement in clinical practice

More data and research will be needed to establish reference ranges in the general population.

Sensitivity and specificity of NfL-based predictions, by using larger cohorts of controls, and taking into account relevant comorbidities and treatment effects.

Assay protocols will need to be standardized and validity of the assay will need to be tested across different populations.

*Neurology. 2018 Sep 12*
References