Improved Differential Diagnosis of Alzheimer’s Disease by Integrating ELISA and Mass Spectrometry-Based Cerebrospinal Fluid Biomarkers

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Handling Associate Editor: Henrietta Nielsen

Accepted 14 November 2018

Abstract

Background: Alzheimer’s disease (AD) is diagnosed based on a clinical evaluation as well as analyses of classical biomarkers: Aβ42, total tau (t-tau), and phosphorylated tau (p-tau) in cerebrospinal fluid (CSF). Although the sensitivities and specificities of the classical biomarkers are fairly good for detection of AD, there is still a need to develop novel biochemical markers for early detection of AD.

Objective: We explored if integration of novel proteins with classical biomarkers in CSF can better discriminate AD from non-AD subjects.

Methods: We applied ELISA, mass spectrometry, and multivariate modeling to investigate classical biomarkers and the CSF proteome in subjects (n = 206) with 76 AD patients, 74 mild cognitive impairment (MCI) patients, 11 frontotemporal dementia (FTD) patients, and 45 non-dementia controls. The MCI patients were followed for 4–9 years and 21 of these converted to AD, whereas 53 remained stable.

Results: By combining classical CSF biomarkers with twelve novel markers, the area of the ROC curves (AUROCS) of distinguishing AD and MCI/AD converters from non-AD were 93% and 96%, respectively. The FTDs and non-dementia controls were identified versus all other groups with AUROCS of 96% and 87%, respectively.

1These authors contributed equally to this work.

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Conclusions: Integration of new and classical CSF biomarkers in a model-based approach can improve the identification of AD, FTD, and non-dementia control subjects.

Keywords: Alzheimer’s disease, cerebrospinal fluid, ELISA, mass spectrometry, mild cognitive impairment, proteomics

INTRODUCTION

Currently, Alzheimer’s disease (AD) is diagnosed based on a clinical evaluation with support from imaging techniques as well as analyses of Aβ42, total tau (t-tau), and phosphorylated tau (p-tau) in cerebrospinal fluid (CSF) [1]. The combination of decreased Aβ42, increased t-tau and p-tau is indicative of AD with a sensitivity of 71–95% and a specificity of 44–87% [2, 3]. Importantly, the sensitivity can be even lower at prodromal disease stages, i.e., in patients with an amnestic form of mild cognitive impairment (MCI) who later will convert to AD dementia [4]. Using logistic regression, studies have evaluated the diagnostic accuracy of combining the classical CSF markers with additional proteins such as YKL-40, NFL, and neurogranin [5–8]. However, such composite biomarker profiles for AD have so far not resulted in any diagnostic improvements [8], implying that there is still a need to identify additional biomarkers to further improve the diagnostic accuracy of CSF analysis.

Here, we combined the classical AD CSF biomarkers with mass spectrometry (MS) based shotgun proteomics data to assess if multivariate modelling using sparse partial least squares discriminant analysis (sPLS-DA) could improve the diagnostic accuracy of recognizing AD from healthy controls and from other conditions of cognitive dysfunction.

METHODS

Samples

This study includes CSF samples from 76 AD patients, 74 mild cognitive impairment (MCI) patients, 11 frontotemporal dementia (FTD) patients, and 45 non-dementia controls. The MCI patients were followed for 4–8 years at 6–12 months intervals and eventually diagnosed with AD (MCI/AD converters) (n = 21) or remained at the MCI stage (stable MCI) (n = 53). Samples were collected according to the recommended consensus protocol for CSF collection and biobanking [9]. All patients underwent computed tomography or magnetic resonance imaging scans, caregiver interviews, thorough cognitive assessments, and in some cases regional glucose uptake by positron emission tomography. The diagnosis of probable AD dementia was set according to the National Institute of Neurological and Communicative Disorders and Stroke/Alzheimer’s Disease and Related Disorders Association criteria [10] and the Diagnostic and Statistical Manual of Mental Disorders, Fourth Edition, criteria [11]. MCI was defined according to the International Working Group of MCI [12]. The control subjects were recruited by advertising in the local newspaper and were considered cognitively unimpaired, based on their history and a Mini-Mental State Examination (MMSE) test.

Approximately half of the controls consisted of 87–89-year-old men from the ULSAM cohort, individuals who have been followed since the early 1970s [13].

The Regional Ethical Review Board in Uppsala, Sweden had approved the collection of CSF samples, the conducted research (collection of the samples: 2005-244 and Ö 48-2005; 2005-11-02 and 2006-01-30 as well as the use of samples for the analyses: 2011/044; 2011-02-23). All participants provided their written informed consent before any samples were collected. The main clinical features of the patients are summarized in Table 1 and Supplementary Table 1.

Sample handling and analysis

The detailed description of sample handling and analysis is available in Supplementary Material 1. Briefly, the CSF samples were collected via lumbar puncture into polypropylene tubes. The concentrations of Aβ42, t-tau, and p-tau in CSF were measured using sandwich ELISAs (INNOTEST, Fujirebio, Ghent, Belgium) and procedures accredited by the Swedish Board of Accreditation and Conformity Assessment.

For MS analysis, the samples were first subjected to multiaffinity immunodepletion to deplete the seven most abundant proteins. The proteins in the depleted CSF sample were digested to peptides
using a trypsin/Lys-C mixture, followed by nanoLC-
MS/MS analyses using a 7 T hybrid LTQ FT MS. The
samples were run according to a random order and in
addition, a pool of CSF was run after every eight
biological samples for the purpose of quality con-
trol. Identification and quantification of MS data was
performed using OpenMS [14]. Proteins identified
(q-value<0.05) with five or more significant peptides
(q-value<0.05) were included in downstream analy-
yses. The proteins with less than 20% missing values
in the QC samples and the biological samples and
coefficients of variation (CV) <1 in QC samples were
selected for modeling. The data was transformed to
log2 scale and normalized using cyclic loess nor-
malization on protein level [6]. The levels of MS
based proteins as well as Aβ42, t-tau, p-tau, and
MMSE were adjusted for age and gender using linear
regression [7].

Identification of AD using Aβ42, t-tau, and p-tau
at baseline

To assess diagnostic accuracy of AD based on
Aβ42, t-tau, and p-tau at baseline, we performed clas-
ification of AD (Aβ42 <530 (ng/L) and t-tau>350
(ng/L)) according to Hansson et al. [15]. For compar-
ison, the results of additional cutoffs, as suggested by
Hansson et al. [15], were also included.

Multivariate modelling to diagnose AD using
Aβ42, t-tau, and p-tau at baseline

We evaluated if PLS-DA could improve the accu-
ry of diagnosing AD versus non-AD subjects
(MCIs, FTDs, and non-dementia controls) by tak-
ing combinatorial effects of Aβ42, t-tau, and p-tau
into account. A linear model using PLS-DA [20]
was trained using the three components for predic-
tion. Importantly, for training, MCI/AD converters
and MCI/non-AD converters were regarded as a sin-
gle group, “MCI”. Therefore, the model was not
provided with information on whether the MCI
patients were AD converters or not. A leave-one-
out cross-validation was performed to evaluate the
accuracy by calculating an area of the ROC curve
(AUROC) for AD versus non-AD subjects (stable
MCI, FTD, and controls), MCI/AD converters versus
non-AD subjects (FTD, stable MCI, and controls),
FTD versus non-FTD subjects (AD, MCI/AD con-
verters, stable MCI, and controls) and controls versus
all other groups (cognitively declined patients).

Integrative multivariate statistical analysis

Using sparse PLS-DA (sPLS-DA), we evaluated if
a combination of Aβ42, t-tau, and p-tau levels with
levels of proteins evaluated by MS could improve the
diagnostic performance. The training and evaluation
of the model, was the same as for the PLS-DA model
described above. The variable importance (VIP) of
the four most important variables (proteins) for each of
the four components used for the diagnostic pre-
diction were automatically selected and extracted
from the model using mixOmics [16]. The AUROCs
were compared between the models using DeLong
test.

Univariate statistical analysis

Nonparametric tests (Kruskal-Wallis, KW) were
performed to evaluate group-wise differences of
Aβ42, t-tau, p-tau, and proteins found in the sPLS-DA
analysis. For statistically significant (p<0.05) results
of the KW test, pairwise post hoc comparisons were
performed using Mann-Whitney U Test. A result of
p<0.05 was considered statistically significant.

Correlation analysis

The Spearman’s rank-order correlation was per-
formed to evaluate association between Aβ42, t-tau,
p-tau and proteins found in the sPLS-DA analysis.

Table 1
Main clinical features information of patients and controls. AD, Alzheimer’s disease; MCI, mild cognitive impairment;
FTD, frontotemporal dementia

<table>
<thead>
<tr>
<th></th>
<th>AD</th>
<th>MCI/AD converter</th>
<th>Stable MCI</th>
<th>FTD</th>
<th>CONTROL</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gender (F/M)</td>
<td>47/29</td>
<td>11/10</td>
<td>20/33</td>
<td>4/7</td>
<td>2/3</td>
</tr>
<tr>
<td>Age (Median[range])</td>
<td>72 (54–88)</td>
<td>71 (59–79)</td>
<td>71 (44–81)</td>
<td>66 (50–75)</td>
<td>88 (74–89)</td>
</tr>
<tr>
<td>Aβ42 ng/l (Median[range])</td>
<td>405 (160–1160)</td>
<td>388 (234–645)</td>
<td>756 (320–1500)</td>
<td>720 (350–1180)</td>
<td>676 (337–1343)</td>
</tr>
<tr>
<td>t-tau ng/l (Median[range])</td>
<td>617 (160–1720)</td>
<td>540 (329–1370)</td>
<td>244 (82–742)</td>
<td>281 (200–600)</td>
<td>414 (202–1121)</td>
</tr>
<tr>
<td>p-tau ng/l (Median[range])</td>
<td>82 (28–220)</td>
<td>78 (35–184)</td>
<td>45 (22–97)</td>
<td>44 (24–59)</td>
<td>63 (29–122)</td>
</tr>
</tbody>
</table>

Age (Median[range]) 72 (54–88) 71 (59–79) 71 (44–81) 66 (50–75) 88 (74–89)
Aβ42 ng/l (Median[range]) 405 (160–1160) 388 (234–645) 756 (320–1500) 720 (350–1180) 676 (337–1343)
t-tau ng/l (Median[range]) 617 (160–1720) 540 (329–1370) 244 (82–742) 281 (200–600) 414 (202–1121)
p-tau ng/l (Median[range]) 82 (28–220) 78 (35–184) 45 (22–97) 44 (24–59) 63 (29–122)
A result of $p < 0.05$ was considered statistically significant.

**RESULTS**

The clinical features of the 206 subjects are summarized in Table 1 and Supplementary Table 1. The 76 AD patients were 54–88 years old and 62% were women. The 74 MCI patients were 59–79 years old and 52% were women. During the 4–8 years follow-up period, 21 of the MCI patients were diagnosed with AD. The eleven FTD patients were 50–75 years old and 57% were women. The non-neurological control subjects were 74–89 years old and 18% were women. Upon KW tests there was a significant age difference ($p < 0.001$), as well as of age and gender adjusted MMSE level ($p < 0.001$) between the groups. The pairwise post hoc testing showed that controls were significantly older than AD, MCI/AD converters, and stable MCI and FTDs ($p < 0.001$). The MMSE scores were significantly ($p < 0.001$) lower in AD compared to both MCI/AD converters and stable MCI.

Identification of AD using Aβ$_{42}$, t-tau, and p-tau at baseline

To assess diagnostic accuracy of Aβ$_{42}$, t-tau, and p-tau at baseline, we performed classification of AD, FTD, MCI/AD converters, and non-dementia controls according to Hansson et al. [15]. The cut off levels of Aβ$_{42}$ <530 (ng/L) and t-tau>350 (ng/L) resulted in an accuracy for identification of AD of 72% (55 out of 76) and incipient AD of 71% (15 out 21). Using this cutoff none of the FTD subjects were classified as AD, but 31% (14 out of 45) of the non-dementia controls were falsely classified as AD. The results using alternative cutoffs, as suggested by Hansson et al. [15], are found in Fig. 1 and Supplementary Table 3. However, the regular cut off levels of Aβ$_{42}$ <530 (ng/L) and t-tau>350 (ng/L) showed the best diagnostic performance.
Fig. 2. Comparison of AUROCs between the classical model (ELISA measurements of $\text{A}\beta_42$, t-tau, p-tau) and the integrative model (ELISA measurements of $\text{A}\beta_42$, t-tau, p-tau in combination with MS-based measurements of 12 proteins). AD, Alzheimer’s disease; MCI, mild cognitive impairment; FTD, frontotemporal dementia.

**Multivariate modelling to diagnose AD using $\text{A}\beta_42$, t-tau, and p-tau at baseline**

We evaluated if PLS-DA modelling could improve the accuracy of diagnosing AD and MCI/AD converters whilst also correctly classifying FTD and non-dementia controls (Fig. 2). This resulted in an AUROC of 92% for discriminating AD versus non-AD subjects and 96% for detecting MCI/AD converters ($p < 0.01$). The AUROC for distinguishing FTD versus all other groups was 57% (not statistically significant). The AUROC for recognition of controls versus cognitively declined subjects was 75% ($p < 0.01$).

**Integrative multivariate modeling to identify incipient AD**

Next, we evaluated if a combination of $\text{A}\beta_42$, t-tau, and p-tau levels with MS based protein measurements could improve the diagnostic accuracy using sPLS-DA. Label free shotgun MS was used to analyze the proteome in all CSF samples. A total of 672 proteins were identified and quantified. After applying sample coverage and CV cutoffs, 78 proteins remained for downstream analyses.

Using sPLS-DA the AUROC for identifying AD versus non-AD was 93% and the recognition of incipient AD (MCI/AD converters) was 96% versus non-AD. The AUROC for distinguishing FTD versus non-FTD increased to 96% ($p < 0.01$). As for recognition of controls versus all other groups, AUROC increased to 87% ($p < 0.01$) (Fig. 2). Comparing the AUROC for the model on the classical biomarkers to the integrated model, the improvement on distinguishing controls versus others and FTD versus others were statistically significant ($p < 0.005$).

**Disease-associated proteins**

Using sPLS-DA we evaluated the different proteins relative contribution to the model predictions...
and SPARC (VIP = 0.15, Uniprot AC: P09486, Uniprot ID: SPRC). Cystatin-C (VIP = 0.43, Uniprot AC: P01034, Uniprot ID: CYTC) Uniprot AC: P36222, Uniprot ID: CH3L1, also known as YKL-40), P04217, Uniprot ID: A1BG), chitinase-3-like protein 1 (VIP = 0.5, Uniprot AC: Q14624, Uniprot ID: ITIH4), apolipoprotein A-I (VIP = 1.75, Uniprot AC: P02647, Uniprot ID: APOA1), Secretogranin-2, beta-Ala-His dipeptidase, alpha-1B-glycoprotein, chitinase-3-like protein 1, also known as YKL-40), cystatin-C and SPARC (Fig. 3). They were in decreasing order: Aβ42, t-tau, p-tau, cadherin-2, neurosecretory protein VGF, afamin, plasma protease C1 inhibitor, inter-alpha-trypsin inhibitor heavy chain H4, apolipoprotein A-I, Secretogranin-2, beta-Ala-His dipeptidase, alpha-1B-glycoprotein, chitinase-3-like protein 1 (also known as YKL-40), cystatin-C and SPARC.

Univariate statistical testing on the selected proteins

The results of KW and Mann-Whitney tests on age and gender adjusted CSF protein levels are illustrated in Fig. 4. The CSF levels of Aβ42 were lower for AD and MCI/AD converters compared to stable MCI, FTDs, and controls. The levels of t-tau and p-tau were higher in AD and MCI/AD converters compared to stable MCI, FTD and controls. The levels of t-tau and p-tau were lower in FTD compared to controls. The levels of plasma protease C1 inhibitor were higher in FTD compared to controls, AD, MCI/AD converters, and stable MCI. The levels of apolipoprotein A-I were higher in AD, stable MCI, and FTD compared to controls. The levels of beta-Ala-His dipeptidase were lower in AD, stable MCI, and FTD compared to controls. Chitinase-3-like protein 1 (YKL-40) was the only protein with higher levels only in AD and MCI/AD converters compared to controls. The levels of Cystatin-C were lower in FTD compared to controls. Finally, the levels of SPARC were higher in AD, MCI/AD converters, stable MCI, and FTD compared to controls (Supplementary Table 4).

Correlations between CSF biomarkers

Figure 5 shows the correlation matrix between Aβ42, t-tau, and p-tau and the twelve MS-based proteins. Aβ42 showed a statistically significant correlation to t-tau, p-tau, VGF, and YKL-40. Total-tau showed a statistically significant correlation to all the analyzed biomarkers except plasma protease C1 inhibitor and cystatin-C. Levels of p-tau were also found to be correlated to the levels of all the markers, except for plasma protease C1 inhibitor. In general, the levels of most proteins showed an inter-correlation, with the exception of plasma protease C1 inhibitor, YKL-40, and SPARC that were found to correlate less frequently to other proteins.

DISCUSSION

The use of CSF biomarkers to support the diagnosis of AD has become gradually more accepted and is today broadly used at memory disorder units in many countries. Although the sensitivities and specificities of the ELISA-based measures of Aβ42, t-tau, and p-tau are fairly good there is still a need to develop novel biochemical markers. In this study, we assessed the CSF levels of the classical AD biomarkers but also applied mass spectrometry to identify additional CSF proteins that were evaluated individually and in combination with the ELISA-based markers. In addition to AD and MCI patients, FTD patients and healthy controls were represented among the 206 subjects included.

By only evaluating the combination of CSF t-tau and Aβ42 we found, in agreement with others [8, 15], that AD and MCI/AD converters could be recognized...
Fig. 4. Scatterplots of CSF levels of the analyzed proteins. The levels were compared between the different groups with nonparametric statistical testing. A: Aβ42, B: t-tau, C: p-tau, D: cadherin-2 (Uniprot AC: P19022, Uniprot ID: CADH2), E: neurosecretory protein VGF (Uniprot AC: P15240, Uniprot ID: VGF), F: afamin (Uniprot AC: P43652, Uniprot ID: AFAM), G: plasma protease C1 inhibitor (Uniprot AC: P05155, Uniprot ID: IC1), H: apolipoprotein A-I (Uniprot AC: P02647, Uniprot ID: APOA1), I: beta-Ala-His dipeptidase (Uniprot AC: Q96KN2, Uniprot ID: CNDP1), J: chitinase-3-like protein 1 (Uniprot AC: P36222, Uniprot ID: CH3L1, also known as YKL-40), K: cystatin-C (Uniprot AC: P01034, Uniprot ID: CYTC), L: SPARC (Uniprot AC: P09486, Uniprot ID: SPRC). AD, Alzheimer’s disease; MCI, mild cognitive impairment; FTD, frontotemporal dementia. p-value: ****0–0.0001, **0.0001–0.001, *0.001–0.01, *0.01–0.05.
Fig. 5. Rank based correlations of the selected proteins. Aβ$_{42}$, t-tau, p-tau, cadherin-2 (Uniprot AC: P19022, Uniprot ID: CADH2), neurosecretory protein VGF (Uniprot AC: O15240, Uniprot ID: VGF), afamin (Uniprot AC: P43652, Uniprot ID: AFAM), plasma protease C1 inhibitor (Uniprot AC: P05155, Uniprot ID: IC1), inter-alpha-trypsin inhibitor heavy chain H4 (Uniprot AC: Q14624, Uniprot ID: ITIH4), apolipoprotein A-I (Uniprot AC: P02647, Uniprot ID: APOA1), Secretogranin-2 (Uniprot AC: P13521, Uniprot ID: SCG2), beta-Ala-His dipeptidase (Uniprot AC: Q96KN2, Uniprot ID: CNDP1), alpha-1B-glycoprotein (Uniprot AC: P04217, Uniprot ID: A1BG), chitinase-3-like protein 1 (Uniprot AC: P36222, Uniprot ID: CH3L1, also known as YKL-40), cystatin-C (Uniprot AC: P01034, Uniprot ID: CYTC) and SPARC (Uniprot AC: P09486, Uniprot ID: SPRC). p-value: ****0–0.0001, ***0.0001–0.001, **0.001–0.01, *0.01–0.05.

with an acceptable accuracy (72% and 71% correct classification), although these markers showed a tendency to diagnose non-dementia controls as AD subjects. Allowing for combinatorial effects of Aβ$_{42}$, t-tau, p-tau, using PLS-DA, improved the accuracy of identifying AD and MCI/AD converters, although the accuracies for recognizing FTDs and non-dementia controls remained low. This means that these markers play a very limited role as a diagnostic test for objectively diagnosing AD, but they serve a purpose when AD is suspected and other diseases have been excluded. Therefore, complementary information is necessary for development of a test based diagnostic system.

When combining the classical biomarkers with MS-based markers the diagnostic accuracy of distinguishing AD (including MCI/AD converters) from patients with other cognitive conditions and healthy controls could be improved just marginally (AUROC for AD: 93% compared to 92%; for MCI/AD converters 96% compared to 93%). These findings are in line with the work by Hampel et al. [8], which demonstrated that a combination of additional proteins (NFL, neurogranin, and YKL-40; as measured by ELISA) could improve the identification of AD from controls (AUROC of 86% compared to 84%) and of AD from FTD (AUROC of 82% compared to 80%). However, their combined additional proteins could not result in an accurate identification of FTD versus healthy controls (maximum AUROC of 78%) and more importantly healthy controls from MCI subjects (maximum AUROC of 62%). In our study, the combination of ELISA- and MS-based data substantially improved the identification of FTD as compared to the ELISA-based data alone (AUROC of 96% compared to 57%, versus all other groups). In addition, the non-dementia controls could also be recognized with a much improved accuracy as
compared to the ELISA-based data alone (AUROC of 87% compared to 75%, versus all other groups).

**Multivariate modeling**

The modelling approach of combining CSF Aβ42, t-tau, and p-tau with MS-based markers has several implications. Importantly, the information regarding which MCI subjects that subsequently converted to AD was not provided to the model at the training stage. By excluding this information, we maintained a high level of stringency and avoided bias for early diagnosis of AD by instead letting the model extract early AD pattern from the data. Therefore, the estimated accuracies for AD and MCI/AD converters should rather be viewed as underestimations. Also, despite that our model increased the risk of misclassification (since the classification was done for each group against all other groups instead of in a pair-wise fashion) it could improve the diagnostic accuracy for AD and FTD as well as more accurately recognize non-dementia controls.

**Disease-associated proteins**

To our knowledge this report represents the largest MS based proteomics study on AD CSF published to date. We undertook a very rigorous experimental approach, where we created a CSF QC pool, which was divided into subsamples that were treated like all other samples, i.e. by depletion, digestion and measurement. This approach allowed us to set an extra stringent cutoff level for the CV of proteins that were included in the final analyses. In addition, we deliberately used a very conservative cutoff for the inclusion of proteins (at least five unique peptides, as compared to the standard procedure of including only two [17]). This conservative approach necessarily leads to an underestimation of the number of proteins that could have been included in the analyses, at the benefit of only including high-abundant proteins displaying low technical variation.

Importantly, the three traditional markers (Aβ42, t-tau, and p-tau) were identified as the top three variables. In addition, the automatic variable selection resulted in the identification of twelve others proteins that were useful for the discrimination of AD, MCI/AD converters, stable MCI, FTD, and non-dementia controls. Nine of these proteins were found to be altered across the groups, according to the KW test.

**Additional AD markers**

The VGF protein was ranked high in the list, indicating its value of being included in a panel of biomarkers for AD differential diagnosis. In agreement with previous reports we found decreased levels of VGF in AD compared to non-dementia controls [18–21] as well as in FTD compared to controls [18, 21]. Moreover, we found that the VGF levels were increased in stable MCI compared to AD, which is also in agreement with recent findings [21]. However, we did not find any difference between MCI/AD converters and stable MCI, contrary to what was recently reported [21].

In agreement with other studies [22–27], we found increased CSF levels of YKL-40 in AD and MCI/AD converters compared to non-dementia controls. Increased levels of YKL-40 in FTD compared to both controls [5] and AD [8] have been previously reported. However, in the current study we did not find any increase in FTD compared to non-dementia controls (p = 0.07), which may be due to the limited number of FTDs included (n = 11). The value of adding YKL-40 in combination with traditional markers is likely to be associated with distinguishing between dementia and non-dementia controls and to some extent between AD and FTD.

**Markers of neurodegeneration and cognitive decline**

The increase in afamin CSF levels in AD, stable MCI, and FTD compared to controls, which correlated strongly to those of apolipoprotein A-I (correlation coefficient of 0.73), have, to our knowledge, not been previously demonstrated. Moreover, the CSF levels of beta-Ala-His dipeptidase were found to be lower in AD, FTD and stable MCI compared to controls.

We found increased CSF levels of apolipoprotein A-I in AD, stable MCI, and FTD compared to non-dementia controls. The results with respect to AD corroborate those of a previous study [28], whereas other studies have failed to replicate those results [29, 30].

SPARC was also found at lower levels in all the patient groups compared to control subjects, indicating its usefulness as a more general biomarker for cognitive decline. The closely related SPARC-like 1 protein has previously been reported as being increased in CSF in AD and MCI compared to controls [31].

Thus, these four proteins seem to have a more general role in neurodegeneration [28, 32–34] and they
probably could serve as biomarkers only when used in combination with other markers.

**FTD markers**

The CSF levels of cadherin-2 were found to be specifically increased in FTD compared to all other groups. The protein was ranked just after the traditional markers, meaning that it had a high value to discriminate the different groups from each other.

We found plasma protease C1 inhibitor to be higher in FTD compared to controls, which contradicts the hypothesis that this protein is AD specific, as previously suggested [35].

For cystatin-C we were not able to detect the previously reported decrease in AD compared to controls [36]. However, the levels were decreased in FTD compared to all other groups, is in accordance with a previous study [37]. Cystatin-C has been implicated as a potential diagnostic marker in both AD, other neurodegenerative diseases and cognitive decline [38, 39]. The CSF levels of cystatin-C showed significant correlation with plasma protease C1 inhibitor and cadherin-2 but with low magnitude, suggesting that these proteins may be complementary to each other and that their value as biomarkers therefore may depend on covariations with other proteins.

Regarding the interpretations of single proteins, it is of importance to emphasize that several markers correlate with each other, but the magnitudes of the correlations were in general small indicating that they indeed are complementary to each other and thus could be used in a combinatorial setting. In line with this, the value from the remaining three proteins were most likely also based on covariations with other proteins, making it harder to interpret their respective values. Among these, inter-alpha-trypsin inhibitor heavy chain H4, secretogranin-2 and alpha-1B-glycoprotein have previously all been associated with AD [36, 40, 41].

**Integrated diagnostics in the clinic**

The molecular events in neurological diseases are complex, emphasizing that no single marker alone can reflect the full pathology. Using a combination of Aβ42, t-tau, and p-tau in CSF is efficient for diagnosing AD and MCI/AD converters, but mainly serves a purpose when other diseases have been excluded. Recent advances in modern technologies have enabled comprehensive measurements of patients suffering from neurological diseases at different molecular levels [42]. Considering the complexity of pathological events, integrating information from multiple sources can therefore result in a more refined tool for diagnostic and prognostic purposes. An example of this is that the incorporation of APOE4 allele information to spectrochemical analysis of CSF can improve the differential diagnosis in AD [43]. We have here demonstrated that adding a limited number of CSF protein measurements in a model-based manner can improve the differential diagnosis. In line with this, we have recently demonstrated that integration of CSF protein, metabolite, and MRI measurements can improve differential diagnosis in multiple sclerosis [44]. Integrative diagnostics holds great potential in future diagnostic assessments yet the challenge is to identify a limited number of markers that holds complementary information and that preferably can be acquired in a clinical setting. The traditional CSF markers (Aβ42, t-tau, and p-tau) play a central role in AD, but by adding proteins to the model we have here demonstrated that this can improve the differential diagnosis to FTD and recognition of non-dementia controls. By inclusion of more disease specific markers, this could be developed into a more general and multipurpose, diagnostic test for neurological diseases that could be used in the clinic. However, translating this into routine healthcare will necessitate rigorous validation process and general acceptance of such a combinatorial test system. This requires more integrative studies to demonstrate and confirm potentials of combinatorial biomarker panels.

**Limitations of the study**

There are a number of limitations to the present study. Firstly, the abundance of MS-based proteins is normalized relative intensities and are thus not representing their respective absolute concentrations. In contrast to Aβ42, t-tau, and p-tau that were measured by a sandwich ELISA. Secondly, the sample size of our cohort was small (especially the FTD group) and all subjects were recruited at one single center. The findings have not been validated in an independent cohort. Therefore, the results might not be applicable to the general population. In addition, there was an age and gender bias, as our non-dementia controls were mostly men and older than the cognitively declined subjects. We controlled for age and gender in our statistical analyses, but an impact from these factors cannot be entirely ruled out. Lastly, the model fitting and calculation of AUROC were based on the latest known health status of the subjects. Therefore,
classification of controls as, e.g., AD or MCI/AD converters can be both due to true misclassification or the subjects having presymptomatic AD.

Conclusions

By combining the ELISA-based classical AD CSF biomarkers with a set of protein markers identified by MS, a marginally improvement of the diagnostic accuracy of AD and incipient AD at the MCI stage could be achieved. FTD patients could be distinguished from non-FTD and non-dementia controls could be distinguished from cognitively declined subjects with significantly enhanced precision. Our findings suggest that incorporating new CSF biomarkers into the currently adopted diagnostic test can further improve the differential diagnosis of AD and concludes that integrative diagnostics holds great potential in future diagnostic assessments.

Further studies are needed to investigate the generalizability of our results.

ACKNOWLEDGMENTS

This research was supported by Uppsala Berzelii Technology Centre for Neurodiagnostics with financing from the Swedish Governmental Agency for Innovation Systems, Swedish Alzheimer’s foundation, Gun och Bertil Stohnes stiftelse, Geriatriiska fonden, Åke Wiberg Foundation and Gamla Tjänarinnor Foundation. The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

Authors’ disclosures available online (https://www.j-alz.com/manuscript-disclosures/18-0855r2).

SUPPLEMENTARY MATERIAL

The supplementary material is available in the electronic version of this article: http://dx.doi.org/10.3233/JAD-180776.

REFERENCES


