

Machine Learning Selection of Most Predictive Brain Proteins Suggests Role of Sugar Metabolism in Alzheimer's Disease

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Abstract.

Background: The complex and not yet fully understood etiology of Alzheimer's disease (AD) shows important proteopathic signs which are unlikely to be linked to a single protein. However, protein subsets from deep proteomic datasets can be useful in stratifying patient risk, identifying stage dependent disease markers, and suggesting possible disease mechanisms.

Objective: The objective was to identify protein subsets that best classify subjects into control, asymptomatic Alzheimer's disease (AsymAD), and AD.

Methods: Data comprised 6 cohorts; 620 subjects; 3,334 proteins. Brain tissue-derived predictive protein subsets for classifying AD, AsymAD, or control were identified and validated with label-free quantification and machine learning.

Results: A 29-protein subset accurately classified AD (AUC=0.94). However, an 88-protein subset best predicted AsymAD (AUC=0.92) or Control (AUC=0.92) from AD (AUC=0.98). AD versus Control: APP, DHX15, NRXN1, PBXIP1, RABEP1, STOM, and VGF. AD versus AsymAD: ALDH1A1, BDH2, C4A, FABP7, GABBR2, GNAI3, PBXIP1, and PRKAR1B. AsymAD versus Control: APP, C4A, DMXL1, EXOC2, PITPNB, RABEP1, and VGF. Additional predictors: DNAJA3, PTBP2, SLC30A9, VAT1L, CROCC, PNP, SNCB, ENPP6, HAPLN2, PSMD4, and CMAS.

Conclusion: Biomarkers were dynamically separable across disease stages. Predictive proteins were significantly enriched to sugar metabolism.

Keywords: Alzheimer's disease, biomarkers, machine learning, metabolism, proteomics, recursive feature elimination

INTRODUCTION

Early elucidation of Alzheimer's disease (AD) is pivotal for constructing clinically impactful treatments. However, the pathophysiology of AD and the driving biochemical changes are not fully understood. Assessment of changes in protein expressions

in the brain may assist in elucidation of multifactorial biochemical changes that lead to AD [1]. Given the complexity and heterogeneity of AD, no single protein is likely to be predictive of all mechanisms or phenotypes which result in AD [2]. Nonetheless, predictive protein models may suggest novel disease mechanisms, improve assessment of patient risk, and signify disease stage-dependent biomarkers [3].

This work identifies protein subsets that differentiate diagnostic labels for AD. AD diagnosis is often based on clinically measured functional cognitive

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decline. AD diagnosis is typically determined using a battery of neuropsychological tests in combination with suggestive imaging, genomic, or other clinical features. Common cognitive tests used in AD diagnosis include the Montreal Cognitive Assessment or Consortium to Establish a Registry for Alzheimer's Disease (CERAD) neuropsychological battery [4]. There is no universal definition of asymptomatic AD (AsymAD). AsymAD is typically characterized by changes in age-adjusted biomarkers, such as increase in amyloid- β and tau in the brain, without overt presence of cognitive decline [5]. Control subjects typically show no overt cognitive losses and no significant change in age-adjusted biomarkers.

In particular, identification of subsets of proteins that better predict and stratify the asymptomatic AD stage is pivotal. Earlier identification of patients likely to transition to AD could enable earlier intervention. The ability to intervene early is likely key to improving outcomes, such as slowing progression or improving symptom-related quality of life. The amyloid- β cascade, tauopathy, and Apolipoprotein E (ApoE) are known aberrant protein signatures in AD [6–9]. However, other proteins may provide earlier clues during asymptomatic changes. For example, metabolomic [10], lipidomic, and inflammation-related proteins have also been suggested to be involved in aging and dementia [11].

The study goal was to determine which proteins in the brain (beyond amyloid- β and phosphorylated tau) are most important for classifying a human subject as either control, AsymAD, or AD. Data consisted of 3,334 brain tissue-derived proteins measured via label-free quantification (LFQ) [3] in six different clinical cohorts. Machine learning classification with recursive feature elimination was used to select the “best” or most predictive proteins.

METHODS

Methods consist of data collection and preprocessing; protein selection using a machine learning algorithm to identify the “best” subset of proteins to predict patient diagnostic classification; validation of the algorithm to accurately classify control, AsymAD, or AD patients using only the identified “best” subset of predictive proteins; and assessment of predictive protein functions. All data preprocessing, machine learning, and analysis was performed in Python 3.6.

Patient diagnostic class labels

Note that the patient diagnostic labels (Control, AsymAD, AD) were inherited from previously published work. Briefly, according to the definitions outlined by Johnson et al. [3], the neuropathological diagnostic classes were determined using CERAD criteria to quantify neuritic plaque distribution and Braak staging to quantify extent of neurofibrillary tangle pathology.

Data used for protein biomarker identification

Six public data sets were utilized [3]: Baltimore Longitudinal Study of Aging (BLSA) [12], Banner Sun Health Research Institute (Banner) [13], Mount Sinai School of Medicine Brain Bank (MSSB) [2], Adult Changes in Thought Study (ACT), Mayo Clinic Brain Bank and University of Pennsylvania School of Medicine Brain Bank. Four data sets ($n=419$ subjects) were utilized for initial model construction and “best” protein selection: BLSA, Banner, ACT, and MSSB. Two data sets ($n=201$ subjects) were used to independently validate the ability of the selected best protein subset to classify the diagnostic label of subjects: Mayo and UPenn cohorts. For all cohorts except Mayo, the tissue was taken from the dorso-lateral prefrontal cortex. For the Mayo cohort, the tissue was taken from the temporal cortex. As shown in Fig. 1a as part of data preparation, missing values were imputed using the k-nearest neighbor (kNN). The optimal number of neighbors for imputation of missing values was determined to be 20 (Supplementary Figure 1). Figure 1b shows the number of subjects and quantified proteins for each cohort. Supplementary Figure 2 illustrates the overall distribution of amyloid- β , tau, APP, CERAD score, and Braak in the data sets used for protein selection. Because amyloid- β and tau were utilized to determine the class labels [3] in the original data sets, tau and amyloid- β levels are not explicitly utilized as part of the protein identification and selection process here. Inclusion of amyloid- β and tau would have resulted in a circular analysis that confounded results. However, their pathways are indirectly represented via upstream biomarkers like APP.

Protein selection using machine learning

As shown in the protein selection row of Fig. 1a, proteins from the selection cohort (data from BLSA, Banner, ACT, and MSSB) were selected using a

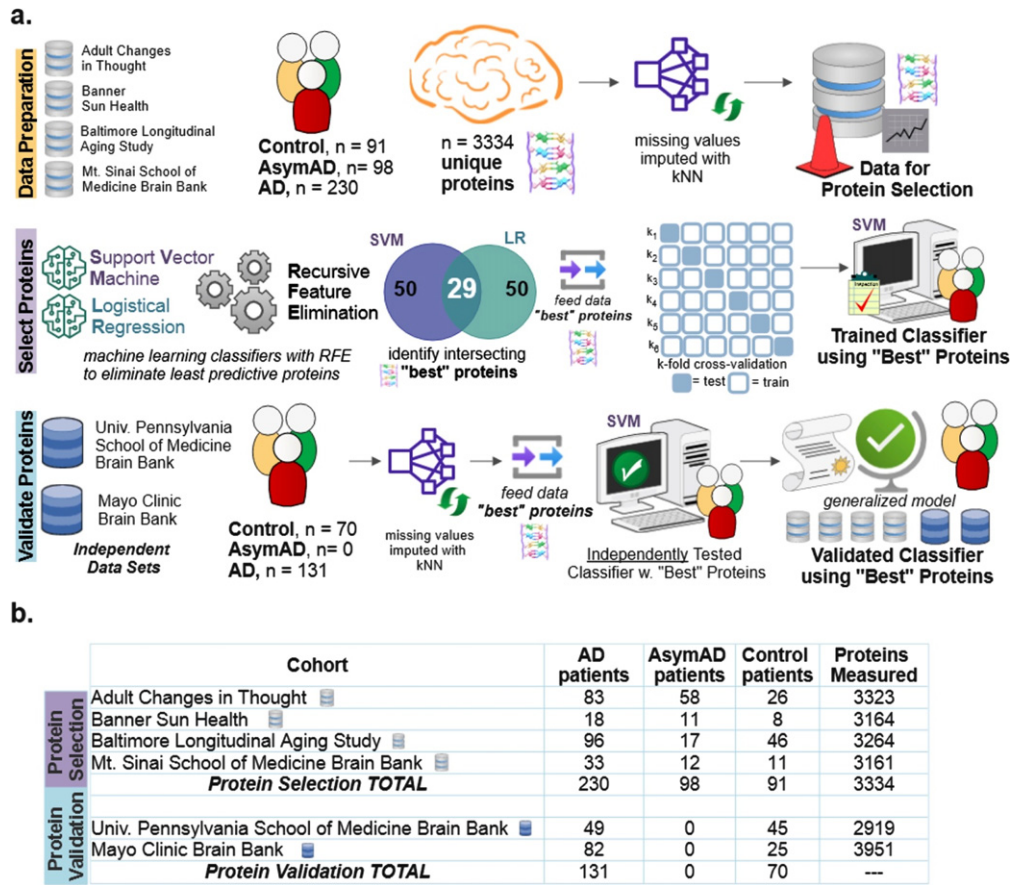


Fig. 1. Diagram explaining the data and machine learning pipeline to identify a subset of "best" predictive protein biomarkers to accurately classify Alzheimer's disease (AD), asymptomatic Alzheimer's disease (AsymAD), or Control. a) Machine learning pipeline consisted of data preparation, protein selection, and model validation with selected proteins. Data was prepared by aggregating four cohorts ($n = 419$ subjects, $n = 3,334$ unique proteins) and imputing missing values using k-nearest neighbor algorithm. The most predictive proteins were selected using recursive feature elimination (RFE) to construct and train a support vector machine (SVM) classifier that can predict diagnosis using only the selected "best" proteins ($n = 29$ or $n = 88$ best predictive proteins). Finally, the developed classifier model was independently validated using 2 additional cohorts ($n = 201$ subjects) to ensure the model's performance generalizes to new data. b) Details of six data cohorts used in protein selection (4 cohorts) and validation (2 cohorts), including sample sizes.

combination of classification algorithms with recursive feature elimination (RFE). RFE is a feature selection algorithm which recursively eliminates less important data features until a pre-defined number of features remain in the dataset. In this study, the "features" are the measured proteins. This iterative procedure is an instance of backward selection [14]. RFE [14] is used to determine the most predictive proteins for successful classification. The resultant predictive protein subset was then used to classify each subject as either control, AsymAD, or AD.

RFE is a wrapper-based feature selection algorithm where recursive rounds of elimination are used to determine the subset of proteins that best predict patient diagnostic classification. The final set of

selected predictive proteins is, in part, sensitive to the classification method. Thus, two popular linear classification methods were independently used with RFE in the scikit-learn package of Python: support vector machine (SVM) and logistic regression (LR), both with linear kernels [15]. The two classifiers, SVM and LR, separately select a specified number of most predictive proteins equal to the RFE criterion. The RFE criterion is the number of proteins the algorithm is allowed to retain. Note that other classifiers were also tried in place or in combination with SVM and LR. However, the intersection of proteins selected by SVM and LR was most consistent and accurate; hence, all results shown utilized this method.

Proteins are selected based on their superior classification ability as quantitatively measured by the area under the precision-recall curve (AUPRC). The intersecting most predictive proteins become the “best proteins”. The RFE algorithm assessed RFE criteria ranging from 10 to 150 proteins. For example, the Venn diagram of Fig. 1a for protein selection illustrates that an RFE criterion of 50 for SVM and LR resulted in an intersecting set of 29 best proteins. Upon completion of protein selection using RFE, a new SVM classifier is constructed, trained, and validated to classify diagnosis (AD, control, AsymAD) using only the selected best proteins. With three classes (control, AsymAD, AD), a one versus rest approach was utilized (AD versus NonAD, AsymAD versus non-AsymAD, control versus non-control).

Note that alternative methods to RFE to identify the most predictive proteins were considered and tried on LFQ as well as held out data: penalized lasso (Supplementary Figure 5), random forest feature importance (Supplementary Figure 6), and statistical differential protein expression using the f-statistic (Supplementary Figure 7). Also, random forests were coupled with RFE to have a more stringent selection criterion - including a protein only when it is selected by three algorithms: SVM, logistic regression, and random forests (Supplementary Figure 8). Performance comparison to neural network, which played no role in feature selection, is also shown in Supplementary Figures 5–8. In some cases, the alternate methods shown in the supplementary figures performed marginally better on the UPenn dataset, which has only binary labels (Control/AD). In all cases, RFE chosen proteins performed substantially better on the LFQ dataset which has more samples ($n=419$), classes (Control/AsymAD/AD), and comprised 4 different datasets (ACT, MSSB, Banner, BLSA) (Fig. 1). Because of its superior multi-class performance and generalizability, the RFE-based primary method shown in Fig. 1a was used to produce all results shown in the main article.

Validation of “best” protein subsets to classify diagnosis

As shown in the Protein Validation row of Fig. 1a, the trained SVM classifier was independently tested using validation cohort data (Mayo, UPenn data sets). As part of independent validation, the best set(s) of proteins determined during protein selection with

RFE was used to predict validation cohort diagnostic classes. However, there were a couple of exceptions due to required data harmonization. In the Mayo cohort, one of the “best” proteins was not quantified (CROCC|Q5TZA2) and a different protein isoform was quantified for APP; APP|A0A0A0MRG2 was included for Mayo, instead of APP|E9PG40). Similarly, for the UPenn cohort, two of the “best” proteins were not quantified (C4A|P0C0L4, DMXL1|Q9Y485), and a different isoform was quantified for APP (APP|A0A0A0MRG2 instead of APP|E9PG40).

Confusion matrices illustrate true positives (TP), false positives (FP), true negatives (TN), and false negatives (FN) used to calculate classification performance. Additionally, a precision-recall curve (PRC) is generated to assess aggregate final model performance. PRC assesses the model’s classification accuracy when using only the “best” protein subsets to classify diagnosis. PRC is a plot of precision versus recall. Recall is defined as $[TP / (TP + FN)]$, and precision is defined as $[TP / (TP + FP)]$. Area under the curve (AUC) provides an aggregate measure of performance across all classification thresholds.

A separate unsupervised learning technique, t-stochastic neighbor embedding (t-SNE), was used to assess separability of AD, AsymAD, and Control subjects using only the selected best proteins subsets determined during supervised learning with RFE.

Finally, principal component analysis (PCA), a dimensional reduction technique, was used to explore and validate RFE criteria. The scree plot and elbow method were used to separately verify how many proteins are necessary to explain the preponderance of variance. The elbow approximated the number of intersecting proteins selected during RFE for optimal diagnostic classification.

Analysis of protein function modules

Selected proteins were matched to their protein function using the color modules and algorithms published by Johnson et al. [3]. There are 14 possible functional modules comprising the entire protein data set ($n=3,334$ unique proteins). The percent composition of specific functional modules in the selected “best” protein subsets were compared to the original, full protein set. Significant differences were assessed using two-sided binomial tests at an alpha of 0.05.

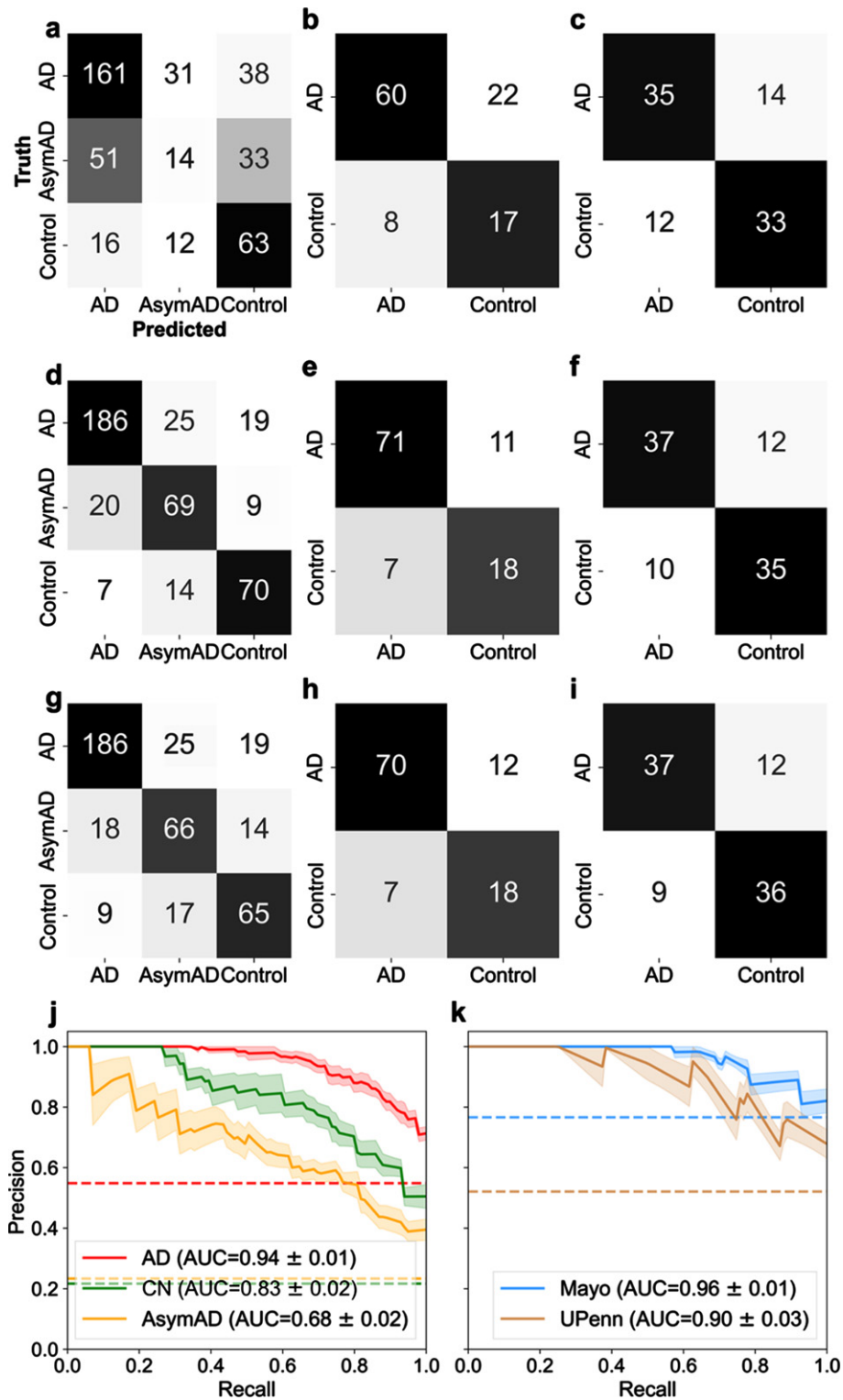


Fig. 2. (Continued)

RESULTS

A machine learning classification and recursive feature elimination process (Fig. 1a) determined which of 3,334 possible clinically measured proteins were most important for classifying control, AsymAD, or AD. RFE was used to identify the proteins that best predicted diagnostic class. Six public data sets were utilized (Fig. 1b). Four data sets ($n=419$ subjects) were for protein selection, which consisted of identifying the “best proteins”. Two data sets ($n=201$ subjects) were used for independent protein validation. Independent validation on unseen data ensured the model was generalizable. Hence, the model can correctly classify the diagnosis of new subjects using only the selected best protein subset(s). An RFE criterion of 50, which resulted in 29 best proteins, was found sufficient to distinguish AD from control. However, an RFE criterion of 150, which resulted in 88 best proteins, was found necessary to optimally distinguish AsymAD from AD.

Classification performance with 29 best proteins

Amyloid precursor protein (APP) is linked to the well-known amyloid- β pathway [16]. It was also selected by RFE as one of the 29 best proteins. Hence, it was important to carefully assess if APP was impacting or biasing classification compared to other proteins. Figure 2 illustrates the classification performance using APP alone (Fig. 2a-c), the selected 29 best proteins (Fig. 2d-f), and APP excluded from the panel of 29 proteins (Fig. 2g-i) for three datasets (LFQ, Mayo, and UPenn). Figure 2d illustrates the performance confusion matrix for the LFQ cohort, Fig. 2e the Mayo validation cohort, and Fig. 2f the UPenn validation cohort, with the selected 29 best proteins. For the LFQ cohort, the model correctly classified 186 of the 230 AD patients (80.87%), while 25 AD patients (10.9%) were misclassified as AsymAD, and 19 AD patients (8.2%) were misclassified as Control. For the Mayo validation cohort, where

proteins were measured in the temporal cortex, 71 AD patients (86.6%) were correctly classified, whereas 11 AD patients were misclassified as Control. For the UPenn validation cohort, where proteins were measured in the dorsolateral prefrontal cortex, 37 AD patients were correctly classified, whereas 12 AD patients (24.5%) were misclassified as Control. Similar trend was seen in Fig. 2g-i when APP was removed from the panel of 29 best proteins and the remaining proteins were used. In summary, the exclusion of APP did not significantly diminish the ability of the remaining 28 proteins to predict diagnosis. This result indicates classification is not dependent on APP alone.

A precision-recall (PR) curve is used to assess aggregate classifier performance. AUC is used to quantify the aggregate classification performance using the selected best protein subset(s). $AUC=1$ is a perfect classifier; thus, an AUC closer to 1 is desirable. Figure 2j illustrates the PR curve and corresponding AUC for the protein selection cohort for AD, control, and AsymAD, respectively, using the selected 29 best proteins. The shaded area represents the standard error, $\pm\sigma$. The 29 best proteins do well in correctly classifying AD ($AUC=0.94 \pm 0.01$; shown in red in Fig. 2j) and control ($AUC=0.83 \pm 0.02$; shown in green in Fig. 2j). However, the best 29 proteins are poor at classifying AsymAD ($AUC=0.68 \pm 0.02$; shown in yellow in Fig. 2j). Figure 2k illustrates the PR curve and corresponding AUC for the independent validation cohorts of Mayo and UPenn, respectively. The AUC for Mayo cohort was 0.96 ± 0.01 whereas the UPenn cohort AUC was 0.90 ± 0.03 . Thus, the model performed equally well in diagnosing unseen AD and Control patients in both independent validation cohorts with the 29-protein subset. The fact validation data originated from different brain regions provides further confidence that the 29-protein subset model is generalizable to other future data sets. The selected proteins do not have a high degree of correlation between them, which supports that the predictive ability does

Fig. 2. Examination of classification performance for Alzheimer's disease (AD), asymptomatic Alzheimer's Disease (AsymAD), and control using $n=29$ best predictive proteins in a one-versus-rest classification setting. Confusion matrices (a-i) illustrate numeric classification results, whereas precision-recall curves (j-k) denote the area under the curve (AUC) with standard error ($\pm\sigma$) to quantify overall classification performance. a-c) Confusion matrices illustrating classification results when using APP alone to classify patient diagnostic class in the (a) LFQ, (b) Mayo, and (c) UPenn datasets. d-f) Confusion matrices illustrating classification results with all 29 “best” or most predictive proteins. g-i) Confusion matrices illustrating results when APP was excluded and the remaining 28 predictive proteins were used for classification of patient diagnosis. APP is widely considered pivotal to the AD etiology. However, these results illustrate APP is not overtly biasing diagnostic classification ability. j) Precision-recall curve for the 3 classes (Control, AsymAD, AD) in the LFQ dataset. k) precision-recall curves for the validation datasets (Mayo and UPenn), which consisted of 2 classes (control/AD). In all cases shown (a-k), an SVM classifier is used with a 6-fold cross-validation strategy, and aggregated results from the test sets are shown.

not rest upon a few proteins in the set (Supplementary Figure 3). Moreover, the unsupervised clustering method, t-SNE, illustrated good separability of the AD, AsymAD, and control classes using the selected subset of 29 “best” proteins (Supplementary Figure 4).

The best 29 proteins are listed in Fig. 3a with their corresponding model coefficient weight as determined from the SVM classification model. Since the one-versus-rest approach for multi-class classification is used, it results in three coefficients for each protein. The three coefficients for every protein correspond to the three diagnostic classes (AD, AsymAD, and Control). The corresponding heatmap illustrates how the selected proteins ($n=29$) drive classification of AD, AsymAD, or Control. Purple represents negative drivers and blue positive drivers. The depth of the hue corresponds to relative magnitude of the coefficient as shown on the heatmap coefficient scale in Fig. 3a. For example, increased APP strongly drives up AD classification, strongly drives down Control, and slightly drives up AsymAD. Similar interpretations can be made for all proteins and their effect on each class.

Figure 3b examines the overlap of the 29 selected proteins in driving diagnostic class (AD, AsymAD, Control). AD and Control, labeled as area 1 on the Venn diagram, share 7 driving proteins: APP, DHX15, NRXN1, PBXIP1, RABEP1, STOM, and VGF. AD and AsymAD, labeled as area 2 on the Venn diagram, share eight driving proteins: ALDH1A1, BDH2, C4A, FABP7, GABBR2, GNAI3, PBXIP1, PKAR1B. AsymAD and Control, labeled as area 3 on the Venn diagram, share seven driving proteins: APP, C4A, DMXL1, EXOC2, PITPNB, RABEP1, and VGF. Note that the color coding of each protein, itself, in Fig. 3 corresponds to function as described in the *Functional Themes in the Selected Proteins* section. The most predictive proteins tend to have opposite signs for coefficient modulation between discriminatory class pairs (AD and control; AD and AsymAD; and AsymAD and Control).

Classification performance with 88 proteins

An RFE of 50, resulting in 29 selected proteins, was sufficient for differentiating AD and Control classes. However, an RFE criterion of 150, resulting in 88 selected proteins, was optimal for differentiating AD and AsymAD classes. Figure 4a and 4b illustrate the PR curve and AUC for each class utilizing the 88 protein subset for predicting diag-

nostic classification. Utilizing the 88-protein subset increased AD and Control classification performance by approximately 4% and 9% respectively compared to the 29-protein subset (Fig. 4a). Utilizing the 88-protein subset increased AsymAD classification performance by approximately 24% (Fig. 4a). In the independent validation cohorts (Fig. 4b), which did not contain any AsymAD patients, the 29-protein subset marginally outperformed the 88-protein subset. In short, AsymAD requires substantially more proteins for accurate predictive classification.

Further exploration of classification performance as a function of RFE criterion

The RFE criterion for protein selection was varied to determine the optimal protein subsets. Again, the RFE criterion determines the number of proteins each classifier (SVM and LR) can select. For a given RFE criterion, the intersection of proteins selected by both SVM and LR become the resultant number of “best” predictive proteins. As described above, RFE=50, resulting in 29 proteins, was sufficient to classify AD versus control. The number of intersecting best predictive proteins for diagnostic classification is not random. Rather, these thresholds are explained by examining dimensional reduction with PCA. Figure 4c illustrates variance explained as a function of number of components. The scree plot approximates minimum components needed to explain the preponderance of variance. The “elbow” of the scree plot denotes the optimal range of components needed to account for the preponderance of variance. Figure 4c shows 29 components (red dot) corresponds to the start of the elbow and 88 components (black dot) corresponds to the end of the elbow. Variance per component beyond the elbow asymptotically approaches zero. Hence, those additional components should not substantively improve model performance. Figure 4d examines the impact of RFE criterion and the resultant number of selected best predictive proteins on diagnostic classification performance. Selecting a RFE criterion greater than 150 (not shown) did not result in increased classification performance.

Functional themes in the selected proteins

The biomarker proteins were mapped to their corresponding modules, which were identified by using the weighted gene correlation network analysis (WGCNA) algorithm. Colors and module number

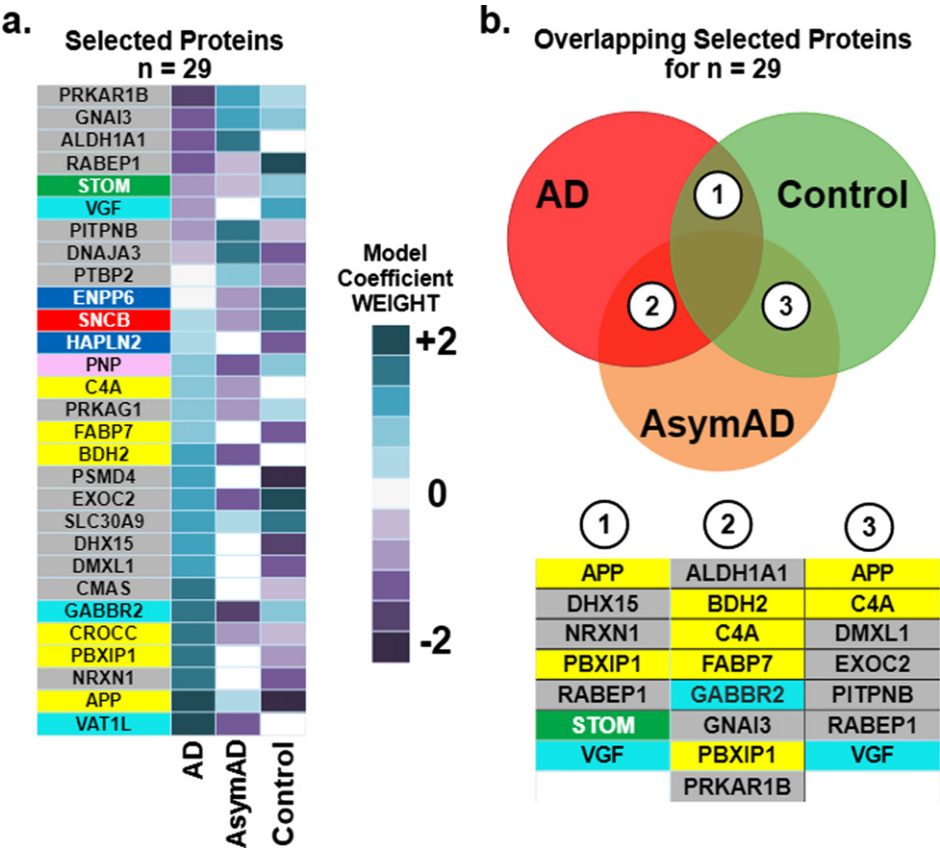


Fig. 3. Driving effects of the selected 29 proteins on the prediction of diagnostic classes of Alzheimer’s (AD), asymptomatic Alzheimer’s (AsymAD), and Control. Note the color of the box around individual listed proteins corresponds to functional modules derived in Johnson et al. [3] and detailed later in Fig. 6. a) The heatmap illustrates the relative magnitude that each protein drives diagnostic class. Driving effects are determined from the coefficients of the SVM. Since the one-versus-rest approach for multi-class classification is used, results encompass three coefficients for each selected protein, one coefficient corresponding to each diagnostic class (AD, AsymAD, Control). b) Key proteins that drive correct prediction in multiple or overlapping classes. The Venn diagram illustrates the overlap in classes: area 1 denotes overlap between AD and Control, area 2 overlap between AD and AsymAD and area 3 between AsymAD and Control. The biomarker panels for each overlap area denote individual predictive proteins shared by overlapping classes.

are used to define the function of the biomarkers [3] and are recapitulated in the first 3 columns of Fig. 5. Functional module frequency (expressed as a percentage) of selected best proteins (for $n=29$ and $n=88$) were compared to the source frequency for the total protein set ($n=3,334$). Change in frequency of selected protein module compared to source frequency is indicative of relative importance of a functional module in predicting diagnostic class (AD, AsymAD, Control). The most enriched module in selected protein sets for both $n=29$ and $n=88$ corresponds to sugar metabolism. Sugar metabolism (M4, yellow) most strongly correlated with AD associated traits (cognition $r=-0.67$, $p=8.5 \times 10^{-23}$; neurofibrillary tangle $r=0.49$, $p=4.7 \times 10^{-27}$; amyloid- β plaque $r=0.46$, $p=1.3 \times 10^{-23}$ and functional sta-

tus $r=0.52$, $p=2.6 \times 10^{-12}$), as reported in [3]. The 29 best proteins are significantly ($p<0.05$) enriched with the sugar metabolism module (M4, yellow). Proteins belonging to sugar metabolism (M4) constituted 5.6% of 3334 total proteins analyzed. However, sugar metabolism proteins constituted 20.7% of the selected best 29 proteins and 13.6% of the selected best 88 proteins (Fig. 5). The remaining functional modules are not significantly different in their representation in either set of selected best proteins. Figure 6 lists the individually selected 29 best proteins ($n=29$) and 88 best proteins ($n=88$) color-coded by functional module. Note the 29-protein set is a subset of the 88-protein set (e.g., the best 29 proteins are all present within the best 88 proteins).

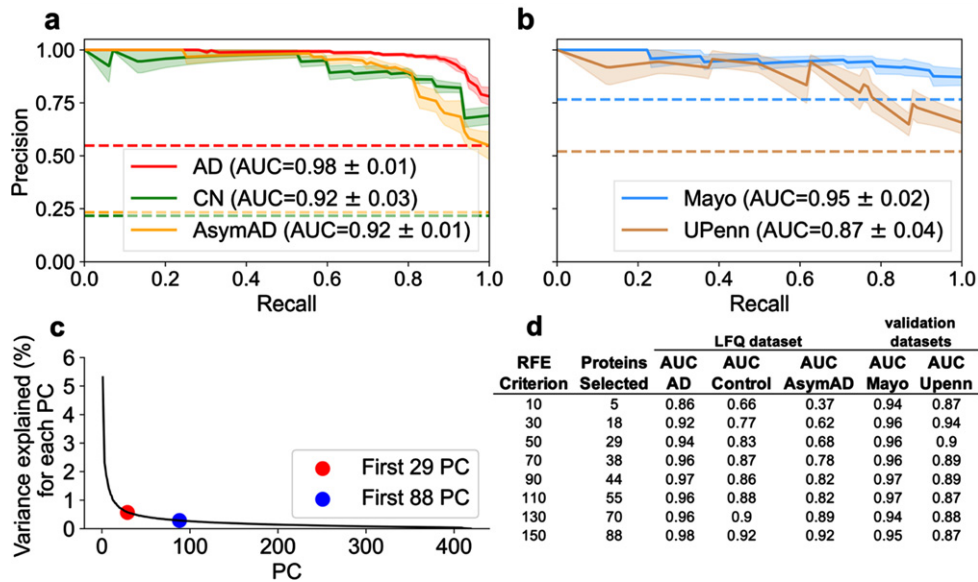


Fig. 4. Examination of optimal number of proteins necessary for superior classification between AD, AsymAD, and Control. A best predictive protein set of 88 ($n = 88$ proteins) was determined to ensure accurate discrimination between AD, AsymAD and Controls in the LFQ dataset. This was based on an assessment of the RFE criterion, principal component analysis (PCA), and evaluation of classifier using area under the curve (AUC) with standard error ($\pm \sigma$). a) AUC of the precision-recall curve for classification of AD, AsymAD, Control using the $n = 88$ best predictive protein set. b) AUC of the precision-recall curve for classification of Control versus AD using the $n = 88$ proteins in the Mayo and UPenn datasets. Of the 88, Mayo dataset had 77 and UPenn dataset had 63 proteins respectively. c) PCA examining variance explained versus number of principal components. Red dot corresponds to the $n = 29$ selected protein subset and blue dot to the $n = 88$ selected protein subset. The “elbow” of the scree plot ends by about 88 principal components. d) Analysis of impact of RFE criterion and resultant number of selected best predictive proteins in the LFQ dataset and the validation datasets using each respective resultant protein subset for classification.

Module	Color	Function	Source Frequency	Selected Frequency (n=29)	Selected Frequency (n=88)
M1	turquoise	synapse	254 (7.6%)	3 (10.3%)	4 (4.5%)
M2	blue	myelin/oligoden	188 (5.6%)	2 (6.9%)	4 (4.5%)
M3	brown	mitochondrial	206 (6.2%)	0 (0%)	1 (1.1%)
M4	yellow	sugar metabolism	186 (5.6%)	6 (20.7%)	12 (13.6%)
M5	green	extracellular matrix	76 (2.3%)	1 (3.4%)	2 (2.3%)
M6	red	cytoskeleton	66 (2.0%)	1 (3.4%)	1 (1.1%)
M7	black	translation/ ribosome	38 (1.1%)	0 (0%)	0 (0%)
M8	pink	blood	46 (1.4%)	1 (3.4%)	3 (3.4%)
M9	magenta	translation/ ribosome	32 (1.0%)	0 (0%)	1 (1.1%)
M10	purple	RNA binding/splicing	45 (1.3%)	0 (0%)	0 (0%)
M11	greenyellow	chaperone/ protein folding	24 (0.7%)	0 (0%)	0 (0%)
M12	tan	unknown	24 (0.7%)	0 (0%)	2 (2.3%)
M13	salmon	unknown	20 (0.6%)	0 (0%)	1 (1.1%)
grey	grey	---	2129 (63.8%)	15 (51.7%)	57 (64.8%)
TOTAL			3334 (100%)	29 (100%)	88 (100%)

Fig. 5. Functional Protein Module of Selected “Best” Predictive Proteins. The functional protein modules are as defined by Johnson et al. [3]. Source frequency is the frequency of the module in the source protein set ($n = 3334$ unique proteins). The selected frequency is for selected best proteins, $n = 29$ or $n = 88$. The M4 yellow module for sugar metabolism is significantly enriched ($p < 0.05$) in selected proteins compared to their frequency in source. Enrichment of sugar metabolism in the selected predictive proteins signifies their importance to diagnostic classification.

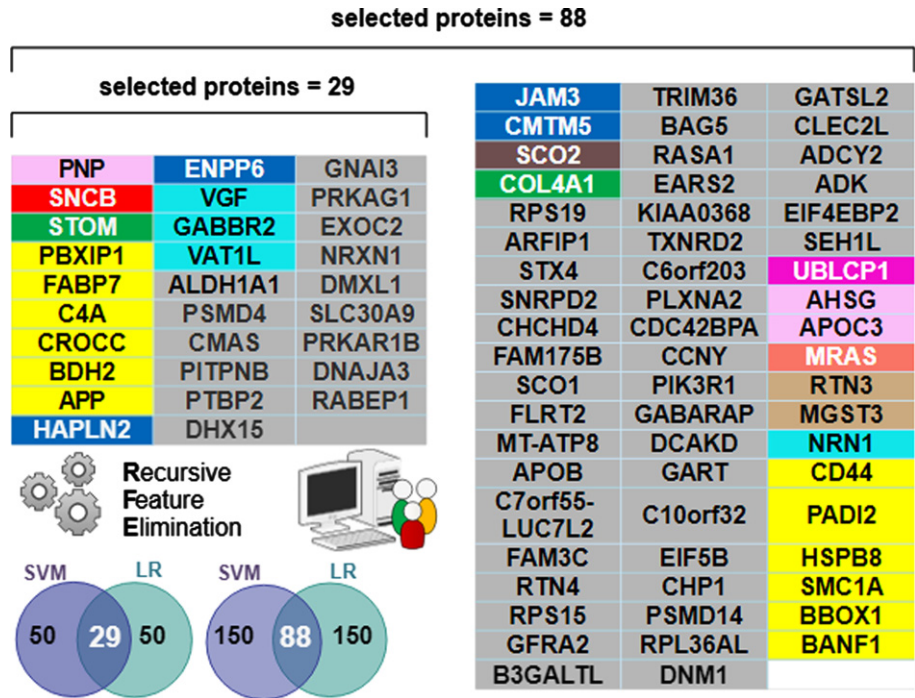


Fig. 6. Individual proteins comprising the selected “best” predictive protein subsets color-coded by functional module. The $n = 29$ subset was sufficient for differentiating AD from Control. However, the $n = 88$ subset was optimal for differentiating AD and AsymAD. Note all 29 proteins in the $n = 29$ selected set are contained within the $n = 88$ selected set. The Venn diagram inset pictorially summarizes the Recursive Feature Elimination (RFE) algorithm used to select the best protein subsets. The intersecting predictive proteins selected by both the support vector machine (SVM) and logistic regression (LR) classifiers during RFE became the “best proteins”. A RFE criterion = 50 resulted in 29 best proteins (e.g., intersection shown on the left Venn inset). A RFE criterion = 150 resulted in 88 best proteins (intersection shown on the right Venn inset).

DISCUSSION

Of the 3,334 proteins, machine learning determined a minimum 29-protein subset necessary to accurately classify AD and Control, but an 88-protein subset was necessary to accurately classify AsymAD. The additional proteins needed for AsymAD classification is likely due to greater complexity and heterogeneity of the AsymAD disease state. The “best” predictive protein subsets ($n = 29$ and $n = 88$) were significantly enriched for sugar metabolism (Fig. 6).

Homeostatic regulatory dynamics key to disease progression

There was relatively little overlap between the predictive proteins that drive control-AsymAD changes and predictive proteins that drive AsymAD-AD changes (see Fig. 3b). This finding indicates an associative relationship to multifactorial dynamic disease progression etiology. In short, the most predictive

proteins dynamically change with disease stage (see Fig. 3a). Whether familial or sporadic AD, different underlying proteomic perturbations may result in multi-scalar system destabilization (e.g., failed homeostasis) with corresponding functional disease phenotypes. Homeostasis is critical for maintaining health, and thus, instabilities often appear in disease [17]. Multifactorial homeostatic instability has been suggested as an underlying propagating mechanism in other neurological pathology, including amyotrophic lateral sclerosis [18], absence epileptic seizures [19], Parkinson’s disease [20], and secondary spinal cord injury [21].

Overlapping proteins for class discrimination

Supplementary Table 1 presents literature details on functions and cited associations with each member of the 29-protein subset. Supplementary Table 1 includes the protein unique ID, brief description of its function and role in AD (if known), and a corresponding reference. Five proteins of the 29-protein subset

overlapped in class discrimination (Fig. 3b): APP, VGF, RABEP1, C4A, PBXIP1. APP (upregulated in AD, AsymAD) was expected given its role in the amyloid cascade [16]. VGF (downregulated in AD) protects against amyloid- β pathology [22]. RABEP1 was key for differentiating Control (upregulated) from AD or AsymAD (downregulated). RABEP1 is tied to longevity and AD [23]. C4A was key for differentiating AsymAD (downregulated) from AD (upregulated) or Control. Increased C4A copy number [24] impacts AD risk and schizophrenia [25]. PBXIP1 was key for differentiating AD (upregulated) from AsymAD (downregulated) or Control. PBXIP1 is cited as altering cell viability and motility through rearrangements of the actin cytoskeleton [26]. Interestingly, many of the 29-proteins are also biomarkers for various non-neural cancers.

Sugar metabolism biomarkers enriched in 29-protein and 88-protein set

Sugar metabolism proteins in both the 29-protein and 88-protein sets (Fig. 6) included: APP, BDH2, C4A, CROCC, FABP7, PBXIP1. Sugar metabolism proteins in the expanded 88-protein subset included CD44, an immune marker associated with AD [27]; PADI2, an age and AD-related marker [28]; BANF1, implicated in aging and progeria [29]; HSPB8, inhibitor of amyloid- β formation [30]; SMC1A, where increased copy number is implicated in epilepsy, AD, and other neurodegenerative diseases [31]; BBOX1, implicated in diabetic kidney disease, lipid metabolic disorders, and schizophrenia [32].

The significantly enriched sugar metabolism module (Fig. 5) supports the recent perspective that asymptomatic and symptomatic AD is characterized by dysregulation of energy metabolism [33, 34]. In short, the presented work supports the hypothesis that sugar metabolism becomes more impacted with disease progression. Insulin resistance in the brain modulates AD inflammatory markers and decreases amyloid clearance [35]. The exact link between AD and type 1 or 2 diabetes is under debate. Nonetheless, poorly controlled blood sugar appears to increase risk of AD [35]. Some researchers have referred to the dysregulation of blood sugar in the brain in AD “type 3 diabetes” [36]. Interconnections between inflammation, metabolism, and protein clearance are further evidence of a multifactorial homeostatic instability contributing to AD progression [3, 10, 34, 37].

The good and the bad of APP

APP is another example where homeostatic instability may play a role in disease progression. For quite some time, APP has been known to be involved in the formation of amyloid- β . A recent genome-wide association study identified APP as a relevant pathway in both familial and sporadic AD [38]. Typically, APP is associated with the formation of toxic soluble amyloid- β oligomers. However, researchers have also suggested that the production of soluble APP alpha (sAPP α) may be a compensatory mechanism to help stave off AD pathology [39]. In particular, amyloid- β monomers have similar neuroprotective properties as sAPP α ; they are neurotrophic and neuroprotective and enhance neurogenesis. Hence, deciphering the possible neuroprotective versus the neurogenic role of APP in AD is an ongoing area of research [40]. The present study cannot confirm or deny the precise causal role of APP as protective, destructive, or a combination of both. The present study’s association-based results do show that APP is an important diagnostic classifier in disambiguating the three stages (Control, AsymAD, AD), as shown in Fig. 3. Nonetheless, the diagnostic classification ability of APP is complex and intertwined with other biomarkers (Fig. 2d-f). When APP was used alone without any other biomarkers, it was not a good classifier (Fig. 2a-c), especially for AsymAD.

Blood-based biomarkers

Biomarkers detected in the blood are preferable for AD risk assessment and early diagnosis [41]. Only one blood-based module protein was selected in the 29-protein subset: PNP, a purine-related metabolite altered early in AD [42]. Two additional blood-based proteins were in the 88-protein subset: AHSG and APOC3. A higher apoE level in high density lipoprotein that lacks apoC3 was associated with better cognitive function [43]. AHSG, a highly glycosylated protein appears downregulated in AD [44].

Assessment of alternatives and limitations

The presented RFE method in Fig. 1, and its corresponding presented results above, were thoroughly vetted and compared to several other statistics-based and machine learning-based model alternatives. The presented method consistently outperformed all other alternative methods and models (see Supplementary Figures 5–8), especially in the mega-LFQ data set

with three classes (Control, AsymAD, and AD). In summary, the presented 29-protein and 88-protein lists for diagnostic classification were quite stable. Relaxing the RFE criterion to include more proteins beyond the selected 88-proteins did not improve classification results (Fig. 4). Nonetheless, no model or method is perfect. While the model is stable, it is fair to expect that a small number of proteins included on the final presented list(s) could be substituted for non-included proteins (e.g., such as similarly co-expressed proteins or proteins from the same functional module). As such, regardless of method, a few proteins that relayed similar, correlated, or mutual information as the selected proteins may not have made the presented final selected proteins list(s). In full transparency, the performance of the protein list generated by each alternative method is shown in Supplementary Figures 5–8. Notably, many of the RFE selected final proteins presented in the main article were recurrently selected by the alternative methods. Finally, any proteins not included in the presented final lists (or even in the input study data) could have their relative importance deduced based on co-expression or their functional modules.

Future directions

The LFQ data covered 3,334 proteins from which the identified “best” biomarker subset is derived. However, the presented method could be extended to future more comprehensive data sets, such as tandem mass tag, to further optimize results. Future addition of larger validation cohorts, especially AsymAD, will ensure model generalizability. Additionally, future inclusion of traits such as gender and race (when available) are important to determine if there are specific feature biases that impact the predictive ability or discriminative expression of proteins. Finally, this work utilized the common 3-class AD staging system: control, AsymAD, or AD. However, it is possible there is a more optimal temporal disease staging system. For example, integrative data machine learning analysis suggested with Alzheimer’s Disease Neuroimaging Initiative data suggested at least four clusters of symptomatic AD patients [45].

Conclusions

Machine learning successfully identified proteins subsets most predictive for classifying AD, AsymAD, and Control subjects. The most predictive proteins subsets comprised < 3% of the 3,334 proteins

assessed. A 29-protein subset accurately classified AD versus Control, but an 88-protein subset was needed to accurately classify AsymAD. The protein subsets resulted in a robust classifier model. The presented model generalized to accurately predict diagnostic labels on unseen data in independent validation cohorts regardless of brain region or minor data set differences. The predictive protein subsets included known important proteins like APP. However, diagnostic classification performance did not hinge upon APP or any single protein or pathway. Finally, the most predictive subsets were significantly enriched in proteins linked to sugar metabolism.

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CONFLICT OF INTEREST

C.M. is an Editorial Board Member of this journal but was not involved in the peer-review process nor had access to any information regarding its peer-review.

DATA AVAILABILITY

All data used in the analysis has been previously published in [3] and is publicly available at <https://www.synapse.org/#!/Synapse:syn20933797/files/>.

SUPPLEMENTARY MATERIAL

The supplementary material is available in the electronic version of this article: <https://dx.doi.org/10.3233/JAD-220683>.

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