

IDENTIFICATION OF DISCRIMINATIVE IMAGING PROTEOMICS ASSOCIATIONS IN ALZHEIMER'S DISEASE VIA A NOVEL SPARSE CORRELATION MODEL

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FOR THE ALZHEIMER'S DISEASE NEUROIMAGING INITIATIVE[†]

Brain imaging and protein expression, from both cerebrospinal fluid and blood plasma, have been found to provide complementary information in predicting the clinical outcomes of Alzheimer's disease (AD). But the underlying associations that contribute to such a complementary relationship have not been previously studied yet. In this work, we will perform an imaging proteomics association analysis to explore how they are related with each other. While traditional association models, such as Sparse Canonical Correlation Analysis (SCCA), can not guarantee the selection of only disease-relevant biomarkers and associations, we propose a novel discriminative SCCA (denoted as DSCCA) model with new penalty terms to account for the disease status information. Given brain imaging, proteomic and diagnostic data, the proposed model can perform a joint association and multi-class discrimination analysis, such that we can not only identify disease-relevant multimodal biomarkers, but also reveal strong associations between them. Based on a real imaging proteomic data set, the empirical results show that DSCCA and traditional SCCA have comparable association performances. But in a further classification analysis, canonical variables of imaging and proteomic data obtained in DSCCA demonstrate much more discrimination power toward multiple pairs of diagnosis groups than those obtained in SCCA.

Keywords: Imaging genomics; Alzheimer's disease; Proteomics; Canonical correlation analysis; Multi-class discrimination.

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1. Introduction

Alzheimer's disease (AD) has been well known as one of the most common brain dementia, a major neurodegenerative disorder that has been characterized by gradual memory loss and brain behavior impairment. According to the latest report,¹ more than 5 million Americans are living with Alzheimer's and it has been officially listed as the 6th leading cause of death. Also, due to the significant decline of self-care capabilities during disease, it is not only the patients who suffer, but also the family members, friends, communities and the whole society considering the time-consuming daily care and high health care expenditures needed. In the past decade, deaths attributed to Alzheimer's disease has increased 68 percent, while deaths attributed to the number one cause, heart disease, has decreased 16 percent. And all of these situations will continue to deteriorate as the population ages during the next several decades. To prevent such health care crisis, substantial efforts have been made to help cure, slow or stop the progression of the disease.

In the last few years, many efforts have been dedicated to explore whether the combination of multi-modal measures, e.g. brain atrophy measured by magnetic resonance imaging (MRI), hypometabolism measured by functional imaging and quantification of proteins, can better predict the clinical outcomes of AD, such as disease status and cognitive outcomes.¹⁹ In many of these works, it has been found that brain imaging and protein expression, from both cerebrospinal fluid (CSF) and blood plasma, hold some complementary information.^{12,18} But how they are related with each other still remains elusive.

In this work, we will explore the relationships between brain imaging and protein expression using bi-multivariate association models. Sparse Canonical Correlation Analysis (SCCA)^{11,16} is a typical example that has been widely used for associative analysis in both real^{8,15} and simulated³ -omics data sets.^{2,11,17} But it can not guarantee the selection of disease-relevant biomarkers and therefore the associations generated in SCCA are not necessarily related to a specific disease either, unless the input features are already prefiltered disease-related biomarkers.⁵ On the other hand, most existing SCCA algorithms use the soft threshold strategy for solving the Lasso^{11,16} regularization terms, which assumes the independence structure of data features. Unfortunately, this independence assumption does not hold in neither imaging nor proteomics data, and will inevitably limit the capability of yielding optimal solutions.

To overcome these limitations, we propose a novel discriminative SCCA (DSCCA) model, coupled with a new algorithm to eliminate the independence assumption, to explore the imaging and proteomic associations. Given imaging, proteomic and diagnostic data, the proposed model can perform a joint association and multi-class discrimination analysis. As such, we can not only identify disease-relevant multimodal biomarkers, but also reveal strong association between them. We perform an empirical comparison between the proposed DSCCA algorithm and a widely used SCCA implementation in the PMA software package (<http://cran.r-project.org/web/packages/PMA/>).¹⁶ The results show that DSCCA and SCCA have comparable association performances. But in a further classification analysis, canonical variables of imaging and proteomic data obtained in DSCCA demonstrate much more discrimination power toward diagnosis groups than those obtained in SCCA.

2. Discriminative SCCA (DSCCA)

Throughout this section, we denote vectors as boldface lowercase letters and matrices as boldface uppercase ones. For a given matrix $\mathbf{M} = (m_{ij})$, we denote its i -th row and j -th column to \mathbf{m}^i and \mathbf{m}_j respectively. Let $\mathbf{X} = \{\mathbf{x}_1, \dots, \mathbf{x}_n\} \subseteq \mathfrak{R}^p$ be the imaging data and $\mathbf{Y} = \{\mathbf{y}_1, \dots, \mathbf{y}_n\} \subseteq \mathfrak{R}^q$ be the protein data, where n is the number of participants, p and q are the number of brain regions and proteins respectively.

Canonical correlation analysis (CCA) is a bi-multivariate method that explores the linear transformations of variables \mathbf{X} and \mathbf{Y} to achieve the maximal correlation between $\mathbf{X}\mathbf{u}$ and $\mathbf{Y}\mathbf{v}$, which can be formulated as:

$$\max_{\mathbf{u}, \mathbf{v}} \mathbf{u}^T \mathbf{X}^T \mathbf{Y} \mathbf{v} \quad s.t. \quad \mathbf{u}^T \mathbf{X}^T \mathbf{X} \mathbf{u} = 1, \mathbf{v}^T \mathbf{Y}^T \mathbf{Y} \mathbf{v} = 1 \quad (1)$$

where \mathbf{u} and \mathbf{v} are canonical loadings or weights, reflecting the significance of each feature in identified associations.

However, the power of CCA in biomedical applications is quite limited due to 1) its requirement on the relatively large number of observations n which is expected to exceed the combined dimension of \mathbf{X} and \mathbf{Y} , and 2) its nonsparse outputs \mathbf{u} and \mathbf{v} which make the ultimate pattern hard to interpret. To address this concerns, sparse CCA (SCCA) method was later proposed, where two penalty terms on both weight vectors $P_1(\mathbf{u}) \leq c_1$ and $P_2(\mathbf{v}) \leq c_2$ were introduced to help generate sparse results.

A widely used SCCA implementation, PMA package,¹⁶ applied L_1 norm penalty for both P_1 and P_2 . But without diagnosis information, its capability in identifying disease-relevant biomarkers is quite limited. Thus the ultimate association relationships are not necessarily related to a specific disease either. Another limitation of PMA is that it takes the soft threshold strategy in the solution, which requires the input data to have an linear independence design $\mathbf{X}^T \mathbf{X} = \mathbf{I}$ and $\mathbf{Y}^T \mathbf{Y} = \mathbf{I}$ (see Section 10 in¹⁴). Unfortunately, this independence assumption does not hold in both imaging and proteomics data (e.g., correlated voxels in an ROI, correlated protein expressions), and will inevitably limit the capability of identifying meaningful imaging proteomics associations.

To overcome these limitations, we propose a novel discriminative SCCA (denoted as DSCCA) algorithm to not only take into account the diagnosis information but also eliminate the independence assumption. Inspired by the application of locality preserving projection (LPP) in linear discriminative analysis,¹⁰ we add two new constraints as P_1 and P_2 for multi-class discrimination.

$$\begin{aligned} P_1(\mathbf{u}) &= \|\mathbf{u}\|_D = \alpha \mathbf{u}^T \mathbf{X}^T \mathbf{L}_w \mathbf{X} \mathbf{u} - (1 - \alpha) \mathbf{u}^T \mathbf{X}^T \mathbf{L}_b \mathbf{X} \mathbf{u}, \\ P_2(\mathbf{v}) &= \|\mathbf{v}\|_D = \alpha \mathbf{v}^T \mathbf{Y}^T \mathbf{L}_w \mathbf{Y} \mathbf{v} - (1 - \alpha) \mathbf{v}^T \mathbf{Y}^T \mathbf{L}_b \mathbf{Y} \mathbf{v}, \end{aligned} \quad (2)$$

Here, we construct two graphs \mathbf{G}_w and \mathbf{G}_b to account for the diagnosis groups, where each vertex indicates one subject (Fig. 1). In \mathbf{G}_w , only subjects within the same diagnosis group have connections to each other. In other words, we build a complete graph for all the subjects belonging to the same diagnosis group. In \mathbf{G}_b , only subjects from different diagnosis

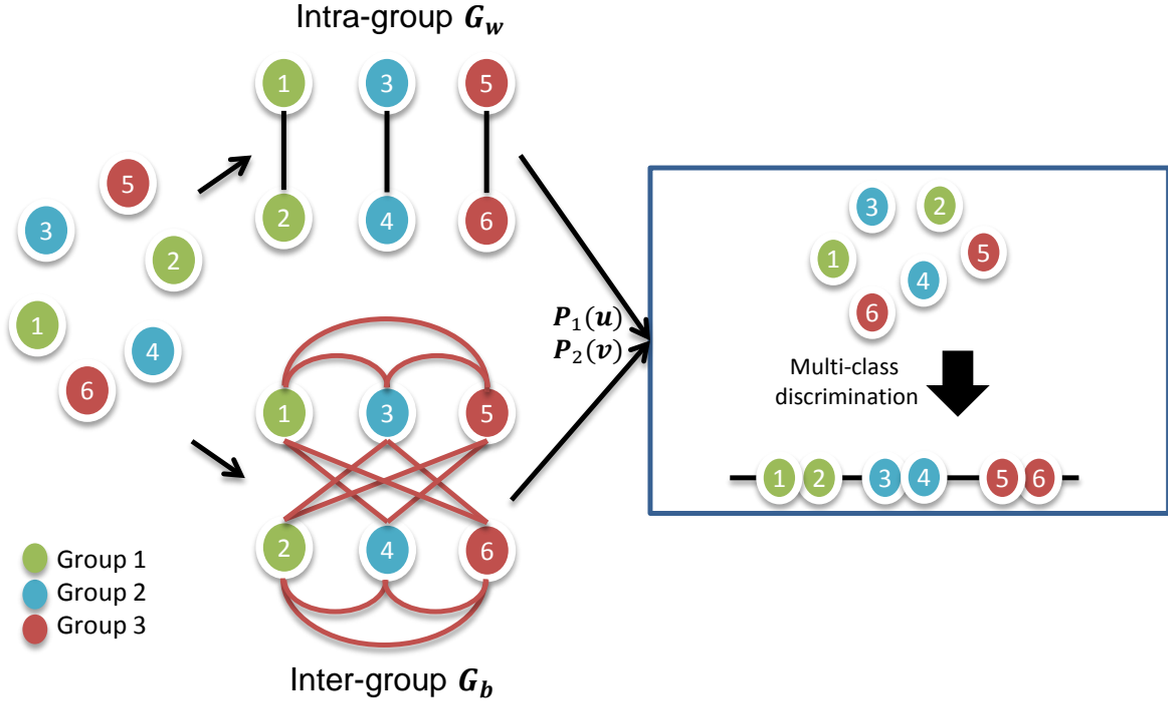


Fig. 1. Illustration of within- and between-group graphs \mathbf{G}_w and \mathbf{G}_b . Each circle indicates one subject and subjects from the same diagnosis group are colored the same.

groups have connections. \mathbf{L}_w and \mathbf{L}_b are the Laplacian graphs of \mathbf{G}_w and \mathbf{G}_b respectively. While the traditional L_1 norm helps ascertain the sparsity of selected imaging and protein biomarkers, the new penalty term $\|\cdot\|_D$ encourages the closeness between subjects within the same diagnosis groups and distance between subjects from different diagnosis groups after projection. α is a trade off parameter that help balance the within- and between-group constraints. Since canonical variables $\mathbf{X}\mathbf{u}$ and $\mathbf{Y}\mathbf{v}$ have the exact same length, we use the same α for both penalties P_1 and P_2 .

The final objective function of DSCCA can be written as follows:

$$\max_{\mathbf{u}, \mathbf{v}} \mathbf{u}^T \mathbf{X}^T \mathbf{Y} \mathbf{v} - \frac{\beta_1}{2} P_1(\mathbf{u}) - \frac{\beta_2}{2} P_2(\mathbf{v}) \quad (3)$$

$$s.t. \quad \mathbf{u}^T \mathbf{X}^T \mathbf{X} \mathbf{u} = 1, \mathbf{v}^T \mathbf{Y}^T \mathbf{Y} \mathbf{v} = 1, \|\mathbf{u}\|_1 \leq c_1, \|\mathbf{v}\|_1 \leq c_2$$

Using Lagrange multipliers, Eq. (3) can be reformulated as follows:

$$\max_{\mathbf{u}, \mathbf{v}} \mathbf{u}^T \mathbf{X}^T \mathbf{Y} \mathbf{v} - \frac{\gamma_1}{2} \|\mathbf{X}\mathbf{u}\|_2^2 - \frac{\gamma_2}{2} \|\mathbf{Y}\mathbf{v}\|_2^2 - \frac{\beta_1}{2} P_1(\mathbf{u}) - \frac{\beta_2}{2} P_2(\mathbf{v}) - \lambda_1 \|\mathbf{u}\|_1 - \lambda_2 \|\mathbf{v}\|_1 \quad (4)$$

Eq. (4) is known as a bi-convex problem, which can be easily solved using an alternating algorithm as discussed in.¹⁶ By fixing \mathbf{u} and \mathbf{v} respectively, we will have the following two minimization problems shown in Eq. (5) and (6).

$$\min_{\mathbf{u}} -\mathbf{u}^T \mathbf{X}^T \mathbf{Y} \mathbf{v} + \frac{\gamma_1}{2} \mathbf{u}^T \mathbf{X}^T \mathbf{X} \mathbf{u} + \frac{\beta_1}{2} P_1(\mathbf{u}) + \lambda_1 \|\mathbf{u}\|_1, \quad (5)$$

$$\min_{\mathbf{v}} -\mathbf{u}^T \mathbf{X}^T \mathbf{Y} \mathbf{v} + \frac{\gamma_2}{2} \mathbf{v}^T \mathbf{Y}^T \mathbf{Y} \mathbf{v} + \frac{\beta_2}{2} P_2(\mathbf{v}) + \lambda_2 \|\mathbf{v}\|_1, \quad (6)$$

Both objective functions can be efficiently solved using the Nesterovs accelerated proximal gradient optimization algorithm.⁹ Algorithm 2.1 summarizes the optimization procedure. The convergence is based on the value changes of the objective function and we use 10^{-6} as stop criteria. Five-fold nested cross-validation was applied to automatically tune the parameters β_1 , β_2 , λ_1 and λ_2 . According to,² the learned pattern and performance are insensitive to γ_1 and γ_2 settings. Therefore in this paper we set both of them to 1 for simplicity. The optimization method used in steps 3 and 4 is similar to that proposed in.⁹

Algorithm 2.1 Discriminative SCCA (DSCCA)

Require:

$$\mathbf{X} = \{\mathbf{x}_1, \dots, \mathbf{x}_n\}, \mathbf{Y} = \{\mathbf{y}_1, \dots, \mathbf{y}_n\}, \mathbf{L}_w \subseteq \mathfrak{R}^{n \times n}, \mathbf{L}_b \subseteq \mathfrak{R}^{n \times n}$$

Ensure:

Canonical vectors \mathbf{u} and \mathbf{v} .

- 1: $t = 1$, Initialize $\mathbf{u}_t \in \mathfrak{R}^{p \times 1}$, $\mathbf{v}_t \in \mathfrak{R}^{q \times 1}$;
 - 2: **while** not converge **do**
 - 3: Solve Eq. (5) using Nesterov's method and obtain \mathbf{u} ;
 - 4: Solve Eq. (6) using Nesterov's method and obtain \mathbf{v} ;
 - 5: Scale \mathbf{u} so that $\mathbf{u}^T \mathbf{u} = 1$
 - 6: Scale \mathbf{v} so that $\mathbf{v}^T \mathbf{v} = 1$
 - 7: $t = t + 1$.
 - 8: **end while**
-

3. Results

3.1. Data and Experimental Setting

The MRI data, quantification of proteins in CSF and blood plasma were downloaded from the Alzheimer's Disease Neuroimaging Initiative (ADNI) database. The primary goal of ADNI has been to test whether serial MRI, PET, other biological markers, and clinical and neuropsychological assessment can be combined to measure the progression of mild cognitive impairment (MCI) and early AD. For up-to-date information, see adni.loni.usc.edu.

We totally extracted 246 subjects with all MRI, CSF and plasma proteomic data available. To balance the diagnostic groups, we randomly removed some mild cognitive impairment (MCI) participants. Finally, 176 subjects (67 AD, 67 MCI and 42 healthy control (HC)), were included in this study (Table 1). For each baseline MRI scan, FreeSurfer (FS) V4 was employed to extract 73 cortical thickness measures and 26 volume measures, as well as to extract the intracranial volume (ICV). CSF and blood plasma samples were evaluated by Rules Based Medicine, Inc. (RBM) proteomic panel and 229 proteomic analytes survived the

quality control process, with 83 from CSF and 146 from plasma. Using the regression weights from HC participants, all the MRI, CSF and blood plasma proteomic measures were pre-adjusted for the baseline age, gender, education, and handedness, with ICV as an additional covariate for MRI only.

Table 1. Participant characteristics

	HC	MCI	AD
Number	67	67	42
Gender(M/F)	38/29	45/22	22/20
Handedness(R/L)	64/3	64/3	38/4
Age(mean \pm std)	75.15 \pm 7.68	74.28 \pm 7.25	75.93 \pm 5.82
Education(mean \pm std)	15.12 \pm 3.01	15.96 \pm 2.92	15.88 \pm 2.77

3.2. Experimental Results

Both DSCCA and PMA were performed on the normalized FS and proteomic measures. To avoid the over-fitting problem, 5-fold nested cross-validation was applied, which also helped to optimally tune the parameters. Table 2 shows 5-fold cross-validation canonical correlation results. It is observed that proposed DSCCA and PMA have comparable performances in identifying imaging proteomic associations, whereas DSCCA is slightly better in performance stability.

Next, we examined the discriminative power of canonical variables $\mathbf{X}\mathbf{u}$ and $\mathbf{Y}\mathbf{v}$ generated by DSCCA and PMA. Area under ROC curve (AUC) was calculated for each single canonical variable of five folds. Both imaging and proteomic canonical variables of PMA and imaging canonical variable of DSCCA were found to have little discrimination power in all HC vs MCI, HC vs AD and MCI vs AD cases. Proteomic canonical variable $\mathbf{Y}\mathbf{v}$ of DSCCA has the best performance, with an averaged AUC around 0.7 for all three cases. Shown in Fig. 2 is an example plot of $\mathbf{X}\mathbf{u}$ against $\mathbf{Y}\mathbf{v}$ in one fold. Dot colors represent different diagnostic groups. Compared to one single canonical variable, we observe that combination of two canonical variables generated in DSCCA demonstrated much more discrimination power than PMA. In Fig. 2(a) three diagnosis groups are all very well separated, whereas in Fig. 2(b) subjects are mixing together.

To further validate our results, a follow up classification analysis was performed using both imaging and proteomic canonical variables as predictors. Canonical loadings learned in the training data set are applied to both training and test data to calculate the training and test canonical variables respectively. The LIBSVM toolbox was employed to implement the SVM using a linear kernel under default settings. Three pair-wise binary classification analyses were performed between HC vs MCI, HC vs AD, and MCI vs AD respectively. Shown in Table. 3 are the classification performance comparison between DSCCA and PMA. The results are very encouraging. Canonical variables of DSCCA significantly outperformed those of PMA in terms of the overall accuracy in almost all the cases. The resulting best prediction rates for HC vs AD (92.1%), HC vs MCI (75.3%) and MCI vs AD (70.3%) were competitive with prior

multi-modal studies,^{6,19} especially considering that it is under default parameter settings.

All five-fold experiments generated similar sparse results in terms of selection of imaging and proteomic markers. Fig. 3 shows the imaging and proteomic markers commonly identified across all folds using DSCCA, where the color represents the weights of corresponding brain regions. Top brain regions identified include entorhinal cortex, amygdala volume, hippocampal volume, etc. (Fig. 3(a)), which are all aligned with previous AD findings.^{12,19} In terms of proteomic markers, expression levels of 12 proteins from CSF and 19 proteins from blood plasma were found to be strongly associated with those brain regions. According to the STRING database (<http://string-db.org/>), these proteins are highly interconnected with each other, as shown in Fig. 3(b). Edges are colored based on the evidence of the connection, such as experimental interaction, co-expression or co-occurrence in the literature. The more edges two proteins have, the more confident their connection will be.

In particular, four proteins, apolipoprotein E (*APOE*), AXL receptor tyrosine kinase (*AXL*), interleukin 6 receptor (*IL6R*) and vascular endothelial growth factor (*VEGF*), were identified in both CSF and blood plasma. *APOE* is the top risk gene of AD. *AXL* is a member of the Tyro3-Axl-Mer (TAM) receptor tyrosine kinase subfamily, which has been previously reported to be involved in Amyloidogenic APP Processing and β -Amyloid Deposition in AD.²⁰ For growth factor VEGF, both its variants and expression changes are found to be associated with AD.^{4,13} *IL6R* is less explored in terms of its relationship with dementia. But in a recent study it was reported to have significant associations with proteins involved in amyloid processing and inflammation.⁷ These findings suggest the existence of certain connections between brain and blood biomarkers. Thus, more accessible fluid biomarkers from blood should have potential to provide extra insights of AD and guidance for future therapeutic intervention activities.

Table 2. Five-fold cross validation canonical correlation results

		f1	f2	f3	f4	f5	mean
DSCCA	Train	0.796	0.670	0.820	0.680	0.636	0.720
	Test	0.424	0.476	0.281	0.392	0.312	0.377
PMA	Train	0.529	0.629	0.505	0.524	0.504	0.538
	Test	0.410	0.095	0.324	0.201	0.460	0.298

4. Discussion

We performed an integrative analysis of brain imaging and protein expression data to jointly identify AD related biomarkers and their associations using a new sparse learning model DSCCA. The overall association performance of DSCCA is better than SCCA. the combination of its two canonical variables are much more powerful in discriminating multiple diagnostic groups simultaneously. Using both imaging and proteomic canonical variables in DSCCA as predictors, we obtained very promising prediction performances: HC vs AD (92.1%), HC vs MCI (75.3%) and MCI vs AD (70.3%), which were competitive with prior multi-modal studies. Since the classification was done under default parameter settings and the sample size is

Table 3. Five-fold cross validation classification performances (%) using canonical variables \mathbf{X}_u and \mathbf{Y}_v . HC vs MCI, MCI vs AD, and HC vs AD are performed as three tasks separately.

	Train			Test			
	HC vs MCI	HC vs AD	MCI vs AD	HC vs MCI	HC vs AD	MCI vs AD	
DSCCA	f1	97.17	100.00	94.19	75.00	91.30	60.87
	f2	86.79	96.51	84.88	85.71	95.65	60.87
	f3	96.23	100.00	94.19	85.71	91.30	86.96
	f4	93.40	95.35	75.58	57.14	100.00	78.26
	f5	72.32	82.61	69.57	72.73	82.35	64.71
	mean	89.18	94.89	83.68	75.26	92.12	70.33
PMA	f1	60.38	77.91	65.12	71.43	86.96	73.91
	f2	66.98	84.88	74.42	71.43	95.65	60.87
	f3	66.04	80.23	63.95	50.00	86.96	60.87
	f4	68.87	80.23	59.30	42.86	82.61	78.26
	f5	65.18	77.17	60.87	31.82	64.71	64.71
	mean	65.49	80.09	64.73	53.51	83.38	67.72

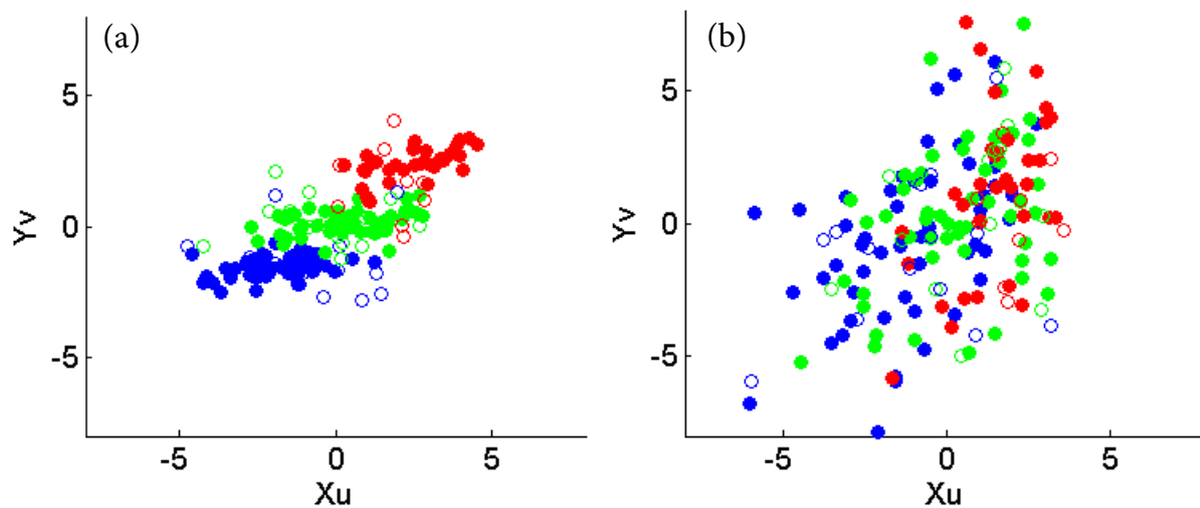


Fig. 2. Plot of canonical variables \mathbf{X}_u and \mathbf{Y}_v . Left: DSCCA; Right: PMA; Red: AD; Green: MCI; Blue: HC; Solid: Training; Circle: Test.

very limited, we expect improved performances with more advanced parameter optimization strategies and/or larger sample sizes.

In real applications, many identified proteomic markers are found to be interconnected, but the underlying mechanisms still warrant further investigation. Replication in independent large samples will be important to confirm these findings. Further pathway enrichment analysis could be performed as a future direction to identify underlying biological pathways of relevant genes and proteins. Considering the ever increasing data volume and diversity in many complex diseases, another potential future topic is to investigate whether DSCCA can help identify valuable complementary information between new -omics features and further improve the classification performance.

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