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Multi-block Analysis of Genomic Data Using Generalized Canonical Correlation Analysis

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Recently, there have been many studies in medicine related to genetic analysis. Many genetic studies have been performed to find genes associated with complex diseases. To find out how genes are related to disease, we need to understand not only the simple relationship of genotypes but also the way they are related to phenotype. Multi-block data, which is a summation form of variable sets, is used for enhancing the analysis of the relationships of different blocks. By identifying relationships through a multi-block data form, we can understand the association between the blocks in comprehending the correlation between them. Several statistical analysis methods have been developed to understand the relationship between multi-block data. In this paper, we will use generalized canonical correlation methodology to analyze multi-block data from the Korean Association Resource project, which has a combination of single nucleotide polymorphism blocks, phenotype blocks, and disease blocks.

Keywords: data interpretation, generalized canonical correlation analysis, genome-wide association study, single nucleotide polymorphisms

Introduction

Human diseases involve complex processes, including the interaction between multiple biological layers, including genetic, epigenetic, and transcriptional regulation [1]. For identifying genes involved in complex human diseases, genome-wide association studies (GWASs) are used by searching for single-nucleotide polymorphisms (SNPs) that occur more frequently in people without the disease. Prior GWASs identified SNPs related to several complex diseases, such as diabetes, heart abnormalities, and Parkinson disease. Researchers hope to find more SNPs associated with chronic diseases through GWASs in the future [2]. Most research has focused solely on the investigation of a single type of genomic data [3].

However, recent advances in genotyping technology have resulted in the large-scale generation of genomic data, and the variety of information collected has also become very diverse. Therefore, it has become very important to apply analytical methods that can fully utilize the given information.

These data need to be analyzed in conjunction with disease variables, such as multiple SNPs and phenotypes. However, since these variables have different properties from each other, multi-block analysis that considers each property is necessary. Furthermore, even though these data have the potential to reveal great insights into the mechanism of disease and to discover novel biomarkers, statistical methods for integrative analysis of multi-block data are only emerging

If there are only two variable blocks, canonical correlation analysis (CCA) can be applied. Research continues to try to analyze DNA-related data based on canonical analysis. Briki and Genest [5] adopted a canonical analysis approach to investigate correlated motions of atoms by molecular dynamics simulation. However, since there are more than two blocks in reality, the extended CCA methods that are applicable to more than two blocks have been less studied.

Hence, research has recently started to shift toward approaches using systematical models in order to integrate and analyze heterogeneous data comprehensively rather than through simple step-wise processes. Among multi-block

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analysis methods, generalized canonical correlation analysis (GCCA) is one of the most representative methods. Nevertheless, many studies have not yet been conducted on the application of multi-block methods to genomic data. Therefore, in the present study, we adopt GCCA to figure out the associations between SNP block, phenotype block, and disease block. We applied Korean Association Resource (KARE) data and analyzed the possibility and limitations of multi-block analysis methods for genomic data.

Methods

Multi-block dataset

We focused on analyzing a multi-block dataset, considering the characteristics of each block. The multi-block dataset is a data type of horizontally concatenating more than two variable blocks. Usually, each block has different properties and forms and can also be partitioned by prior knowledge, but all blocks have the same number of observations. Suppose there are K blocks, and each block has p_k number of variables ($k = 1, \dots, K$). We can express the k-th block X_k as $X_k = [x_1, \dots, x_{pk}]$. The total dataset X can be presented as $[X_1, \dots, X_k, \dots, X_K]$.

As technology advances, this type of data is common in a variety of studies. For instance, in food science, blocks of variables could be physico-chemical measurements, sensory analysis data, and instrumental measurements [6]. Multiomics, the typical multi-block dataset in the medical field, means a new biological analysis approach where the datasets are multiple omics, such as the genome, proteome, transcriptome, epigenome, and microbiome [7-9]. It usually focuses on associations between SNPs and traits, considering varying phenotypes.

Generalized canonical correlation analysis

Among various methods of dealing with a multi-block dataset, GCCA is used in this paper. Since it is extended from CCA, we start with an explanation of CCA. CCA is a method of inferring information from cross-covariance matrices. If

there are two vectors $X = (x_1, \dots, x_n)$ and $Y = (y_1, \dots, y_m)$ of random variables and if there are correlations among the variables, the linear combinations of x_i and y_j to maximize the correlation with each other—termed canonical variables—are found through CCA [10].

GCCA is a way of extending CCA to adapt to more than two sets of random variables after removing dependencies within each set. The basic structure of CCA is to derive a new linear combination of the variables, called canonical variables, constituting each set and to estimate the correlation between canonical variables. In other words, canonical variables summarize the information inherent to the abbreviated set of multivariate data [11]. GCCA can be divided into two methods: using correlations and using covariance. In our paper, we use a method of analysis based on covariance. GCCA based on covariance uses the variance of 'block scores' to compute the residual matrices. For instance, in an X_k variable block, we can denote $a_k = (a_{k1} \ a_{k2} \cdots$ a_{kp_b})' as the coefficients for each variable in X_k block. Therefore, the canonical variables, y_k (k = 1, ···, K), are expressed as:

$$y_1 = X_1 a_1 = a_{11} X_{11} + \dots + a_{1p_1} X_{1p_1}$$

$$\vdots$$

$$y_K = X_K a_K = a_{K1} X_{K1} + \dots + a_{Kp_K} X_{Kp_K}$$

The optimization problem is as follows.

$$\underset{\left\{a_{1}, a_{2} \cdot \cdot \cdot \cdot a_{k}\right\}}{\operatorname{argmax}} \quad \sum_{i, j = 1, i \neq j}^{K} c_{ij} g(cov(X_{i}a_{i}, X_{j}a_{j}))$$

The optimization problem tries to find coefficients of each block a_1 , a_2 , \cdots , a_K that would maximize the weighted summation of the covariance of the two components. The c_{ij} in the equation implies the relationship between variable block X_i and X_j . If they have a relationship, we could assign $c_{ij} = 1$; otherwise, we could assign $c_{ij} = 0$. The function g()

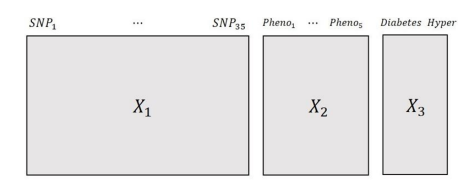


Fig. 1. Construction of multi-block dataset.

can be various functions, such as horst (g(x) = x), centroid (g(x) = |x|), and factorial $(g(x) = x^2)$. Among these methods, we applied horst methods. A design matrix $C = (c_{jk})$ is pre-specified by the user to express the relationships between blocks. The element c_{jk} is equal to 1 if block j and block k are connected and 0 otherwise [12, 13].

Data description: KARE

In this paper, we use data from the KARE project, which was initiated in 2007 to undertake a large-scale genome-wide association analysis among the 10,038 participants of two areas: Anseong and Ansan. It was established as part of the Korean Genome Epidemiology study (KoGES) in 2001, providing genomic and clinical variables for over 260 traits [14].

Among the KARE data, we decided to make three different variable blocks that included information on SNP variables, phenotype variables, and disease variables. Fig. 1 represents the multi-block form of the KARE data, which had three variable blocks, each with different characteristics. The first variable block, X_1 , is a block of SNP variables; the second variable block, X_2 , is a phenotype block that has five phenotype variables related to obesity. The last variable block, X_3 , is a disease block that has information on observational status in diabetes and hypertension.

The first variable block, X1, has information on 35 SNP

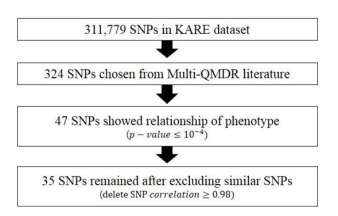


Fig. 2. How to extract 35 single nucleotide polymorphism (SNP) variables. KARE, Korean Association Resource.

variables, and each piece of data was recorded as 0, 1, or 2 according to their genotype. We extracted 35 SNP variables to be included in our analysis according to the specific following steps described in Fig. 2. The original KARE dataset has 311,779 variables, and we regarded 324 SNP variables as our main interest from the literature of Multi-QMDR analysis. The 324 SNP variables showed strong marginal effects in the univariate linear regression models in the paper [15]. From the 324 selected variables, we selected 47 variables that showed a significant relationship with our phenotype variables in the phenotype block. Lastly, we removed extremely similar SNP variables that had a correlation of more than 0.98 with each other in order to clearly see the correlation between variables.

The second variable block, X2, is a block of phenotype variables that have been proven to have a relationship with obesity. The five phenotype variables-suprailiac skinfold, subscapular skinfold, body mass index (BMI), waist-hip ratio, and waist-were selected, and all of them are related to obesity. The third variable block, X₃, is a block of diseases. Two disease variables were made from patients' clinical traits. Participants whose "fasting blood glucose" was higher than 126, "blood glucose/oral glucose tolerance after 120 minutes" was higher than 200, or "who had medication of diabetes" were considered as having diabetes. Participants whose "subscapular skinfold" was over 140, "suprailiac skinfold" was over 90, or "who had medication of hypertension" were considered as having hypertension. Table 1 shows how each group was composed of according to our disease definition. Excluding individuals with missing values among the variables used in this process, the final sample size was 7,389 in the study.

Sex and age were considered as potential covariates which could affect our association analysis; for instance, coronary heart disease (CHD) is more common in men than in women. In addition, the risk of CHD increases with age in both sexes, but the increase is sharper in women [16]. Therefore, we divided the 7,389 observations into four groups based on a median age of 50 years and sex. Group 1 represents below the median age and males, while group 2 represents above the median age and males. Groups 3 and 4

Table 1. Disease frequency according to group

	Group 1		Group 2		Group 3		Group 4	
	Yes	No	Yes	No	Yes	No	Yes	No
Diabetes	119 (6.9)	1,594 (93.1)	151 (10.2)	1,325 (89.8)	74 (3.6)	1,992 (96.4)	206 (9.7)	1,928 (90.3)
Hyper	217 (12.7)	1,496 (87.3)	341 (23.1)	1,135 (76.9)	152 (7.4)	1,914 (92.6)	568 (26.6)	1,566 (73.4)
Total	1,7	13	1,4	476	2,0	066	2,1	134

Values are presented as number (%).

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represent females below and above the median age, respectively. For each group, our data were composed of three different blocks (J=3)—a gene data block, clinical data block, and disease status block. As with many multi-block genomic data, KARE data also have very different characteristics for each block. The gene data block has 35 SNP variables that are discrete and can only have values of 0, 1, or 2. The five phenotype variables of the clinical data block were continuous. The disease status block consisted of only

two dummy variables, indicating the presence or absence of disease.

To check the association between blocks, logistic regression analysis was performed, and the coefficients and p-values are listed in Table 2. Table 2 shows the simple logistic regression of the variables in the phenotype and genotype blocks for diabetes and hypertension.

In Table 2, every variable in the phenotype block was statistically significant with diabetes, all with a p-value less

Table 2. Simple logistic regression coefficients for diseases

DJ I	Dependent variable —	Dial	betes	Hypertension	
Block		Estimate	Pr (> t)	Estimate	Pr (> t)
Phenotype	Suprailiac skinfold	0.021	< 0.001	0.012	0.004
	Subscapular skinfold	0.020	< 0.001	0.002	0.578
	BMI	0.026	< 0.001	0.045	< 0.001
	Waist-hip ratio	0.032	< 0.001	0.070	< 0.001
	Waist	0.033	< 0.001	0.064	< 0.001
Genotype (SNP)	rs221097	0.011	0.511	-0.055	0.015
	rs7583940	0.014	0.105	-0.002	0.858
	rs3103261	0.005	0.723	-0.015	0.486
	rs3856726	-0.045	0.025	0.02	0.489
	rs1849809	-0.003	0.546	-0.003	0.672
	rs7681841	-0.021	0.293	0.008	0.786
	rs17226252	-0.014	0.387	0.009	0.705
	rs1570064	-0.007	0.704	-0.006	0.813
	rs17168600	-0.013	0.284	0.025	0.141
	rs6965746	-0.004	0.347	0.013	0.047
	rs1510447	0.000	0.993	-0.005	0.521
	rs4472504	-0.01	0.394	0.037	0.031
	rs10090537	-0.013	0.535	0.009	0.759
	rs4745034	-0.005	0.689	0.024	0.198
	rs16906215	-0.001	0.957	0.014	0.479
	rs17599042	-0.005	0.652	0.018	0.27
	rs17109716	-0.008	0.527	0.005	0.794
	rs11876341	0.003	0.631	0.017	0.084
	rs601619	0.001	0.942	-0.002	0.953
	rs17248901	-0.024	0.059	0.012	0.515
	rs6561930	-0.037	0.052	0.024	0.381
	rs527248	0.01	0.06	-0.005	0.498
	rs11000212	0.001	0.828	0.001	0.923
	rs9939609	-0.003	0.639	0.024	0.008
	rs10842994	0.002	0.773	0.001	0.853
	rs3782889	-0.011	0.045	-0.016	0.039
	rs12229654	-0.016	0.009	-0.021	0.013
	rs11066280	-0.013	0.019	-0.028	< 0.001
	rs4667458	-0.005	0.571	-0.011	0.418
	rs11933222	-0.005	0.223	0.008	0.205
	rs17092358	0.011	0.032	0.003	0.635
	rs7136259	-0.004	0.341	-0.031	< 0.001
	rs2254613	0.003	0.487	0.028	< 0.001
	rs1378942	-0.011	0.045	-0.037	< 0.001
	rs11131794	0.02	< 0.001	-0.01	0.174

BMI, body mass index; SNP, single nucleotide polymorphism.

than 0.001. However, in the regression with hypertension, the subscapular skinfold variable did not satisfy the significance level, which turned out to be a p-value of 0.578. Other than subscapular skinfold variable, all variables in the phenotype block showed significance of a relationship with hypertension. Therefore, there exists a relationship between each phenotype variable and two diseases, respectively. In contrast to the association between phenotype variables and disease variables, the association between SNP variables and disease variables were revealed only from certain genes. The genes related to diabetes were rs3856726, rs3782889, rs12229654, rs11066280, rs17092358, rs1378942, and rs11131794. The genes related to hypertension were rs221097, rs6965746, rs4472504, rs9939609, rs3782889, rs12229654, rs11066280, rs7136259, rs2254613, and rs1378942. There has been a study of SNPs associated with diabetes, in which the MYL2, C12orf51, and OAS1 genes were found to be significantly associated with 1-hPG, which has been understood as an additional risk factor for type 2 diabetes. Therefore, genes with rs3782889 (MYL2), rs12229654 (MYL2) were proven to have a valid relationship with diabetes [17].

Results

The results of the GCCA were analyzed separately according to each group. In doing the GCCA, we needed to determine how many canonical variables we would have. The generalized canonical correlation results are presented in Table 3. Group 1 had generalized canonical correlations of 0.183, 0.156, and 0.095. The square of the generalized canonical correlations was 0.034, 0.024, and 0.009. The square of the generalized canonical correlation was used for calculating what proportion the canonical variable explains the dataset. For instance, the first canonical variable in group 1 had a square of the generalized canonical correlation value of 0.034 (rounded number is listed in Table 4). The proportion was calculated by using the square of the

Table 3. Correlation between blocks

Group ^a	Canonical variable	SNP-Phenotype	SNP-Disease	Phenotype-Disease
Group 1	First-first	0.204	0.164	0.180
·	Second-second	0.281	0.071	0.071
Group 2	First-first	0.194	0.128	0.186
	Second-second	0.277	0.085	0.034
Group 3	First-first	0.169	0.102	0.250
	Second-second	0.259	0.031	0.032
Group 4	First-first	0.193	0.129	0.188
	Second-second	0.192	0.061	0.069

SNP, single nucleotide polymorphism.

Table 4. Generalized canonical correlation results

Group ^a	Canonical variable	Generalized canonical correlation	Square of generalized canonical correlation	Proportion
Group 1	1	0.183	0.034	50.2
	2	0.156	0.024	36.4
	3	0.095	0.009	13.4
Group 2	1	0.171	0.029	46
	2	0.151	0.023	35.8
	3	0.107	0.012	18.2
Group 3	1	0.178	0.032	52.9
	2	0.133	0.018	29.8
	3	0.101	0.010	17.3
Group 4	1	0.171	0.029	55.9
	2	0.115	0.013	25.2
	3	0.099	0.010	18.9

^aThe groups 1 and 2 represents men group with age below median and age above median, respectively. The groups 3 and 4 represents women group with age below median and age above median, respectively.

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generalized canonical correlation. The first GCCA canonical variable explained 50.2% of the variation, while the second and third variables explained 36.4% and 13.4%, respectively. Therefore, for group 1, we used two canonical variables that were sufficient enough to represent the dataset. The other three groups also required two canonical variables to represent the dataset.

Since we have three blocks, we could make three initial canonical variables. The SNP block's first canonical variable was $U_1 = a_1SNP_1 + \cdots + a_{35}SNP_{35}$. The phenotype block's first canonical variable was $V_1 = b_1SUP + b_2SUB + b_3BMI + b_4WHR + b_5WAIST$. The disease block's first canonical variable was $W_1 = c_1Diabete + c_2Hyper$. The term a_i , b_i , c_i represents the coefficient of each i-th variable in the block. When we draw the first canonical variables U_1 , V_1 , W_1 and the second variables U_2 , V_2 , W_2 , we can interpret the relationship of each block's variable to the other block. In this paper, we first explain the relationship between blocks and then discuss the relationship within the blocks.

Between-block relationship

Once we conducted GCCA in SAS/IML, we could get information on the canonical variables and coefficients for each variable in the block. From the canonical variable value, we calculated the correlation between blocks, because the first and second canonical variables could represent most of the data. Table 3 represents the correlation between each block's canonical variables. With a GCCA containing three blocks, the first correlation among three blocks is obviously high, but it can be different when we consider two pairs. The highest relationship was found in group 1, which had a correlation of 0.281 between the SNP block's second canonical variable and the phenotype block's second canonical variable. It is interesting that the relationship between the SNP block and phenotype block is the strongest, while the relationship between the SNP block and disease block is not that noticeable. This would imply a pathway relationship from SNP to disease, in which phenotype is the medium of the link between them. Another interesting point is the relationship between the first canonical variables is not that strong in the relationship between SNP and phenotype. Except group 4, the other three groups showed a stronger correlation between second canonical variables than first canonical variables. This is thought to be due to the fact that the phenotype spreads in the second axis.

We also visualized the relation between blocks, particularly between the second canonical variables of the SNP and phenotype blocks. In Fig. 3, we illustrate a sample plot in group 3. The plot explains how the sample observation is located in each block by using the first and second canonical variable of each block. The SNP block appeared to be scattered compared with the other groups. The phenotype block's observations are densely populated near the center (0,0), while the disease group's observations only appear at four dots, since they are discrete data that have four possible disease cases. We can regard a sample that is located farther from center (0,0) is more influential for each SNP, phenotype, or disease block.

Within-block relationship

If we looked at each block's variable plot, we can also understand how each variable in the block affects the canonical variable. The variable plot demonstrates how the variables are positioned within block; therefore, the goal is figuring out the internal relationship between variables. The x-axis is how much each variable influences the first canonical variable, and the y-axis is how much each variable influences the second canonical variable. In Fig. 4, there exist four variable plots, each representing how the variable in each block composes its block. The variables of the SNP block were represented as black points, which were categorized into seven different groups, based on where it is related the most.

We can interpret variable plots through making criteria for the x-axis and y-axis. If a variable exists farther away from the origin, the influence would be greater. We will illustrate group 1's case in particular, while the interpretation of the other groups could be made, in addition to the first group's

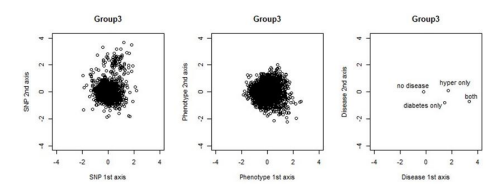


Fig. 3. Sample plot of group 3.

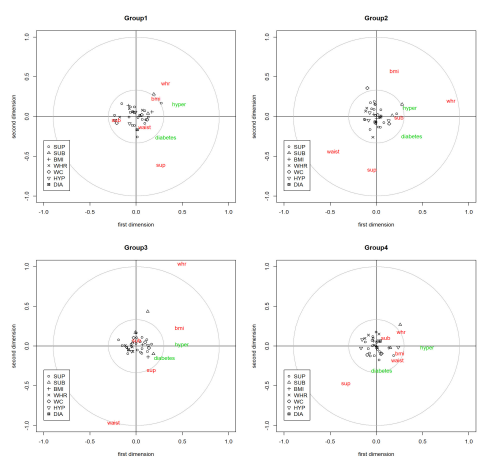


Fig. 4. Variable plot of each group. SUP, suprailiac skinfold; SUB, subscapular skinfold; BMI, body mass index; WHR, waist-hip ratio; WC, waist circumference; HYP, hypertension; DIA, diabetes.

interpretation. In group 1's variable plot in Fig. 4, compared with the SNP variables, phenotype and disease variables showed more distinguishable features. The 35 SNP variables, which were presented as seven symbols according to their category in the graph, had a similar influence on SNP blocks. In the phenotype block, the variable waist-hip ratio and BMI had almost the same angle from the origin, whereas suprailiac skinfold (Sup) was almost in the opposite location in terms of the second components of the phenotype blocks. This means that waist-hip ratio and BMI gave similar traits compared with other phenotypes. When we considered the first-dimension axis, we could observe that subscapular skinfold (Sub) had a different trait from the other phenotype variables. In terms of disease blocks, we could separate people with disease and people without disease in terms of the x-axis, and hypertension and diabetes were also distinguishable with the y-axis criteria.

It is interesting for us to compare how the variable plot in each group was different. In terms of the SNP variable block, the rs527248 SNP variable had a triangular shape, which means the SNP is related to BMI the most compared with the association with other phenotype variables. Among SNP variables, rs527248, a point located outside of small gray

circle in the first quadrant, had the most powerful influence on its canonical variable in all groups. In group 2, the influence of the rs527248 variable was more powerful on the first axis, whereas for groups 1, 3, and 4, there was much influence of the variable on the y-axis.

There are different aspects of the phenotype block's variables from each group. In group 4, there existed extreme difference between the phenotype variables waist-hip ratio and waist. Whereas the waist-hip ratio variable was more than 1 on the y-axis, the waist variable was near –1, which is exactly the opposite location. If we look at the specific coordinates of the waist-hip ratio and waist variables, the influence of waist-hip ratio on the y-axis was 1.038, but the influence on the x-axis was 0.4966. In contrast, the waist variable in group 3 had a stronger influence on the y-axis than on the x-axis.

From our simple logistic regression, we had information that the SNP variable rs4472504 was related to hypertension disease, which had a p-value of 0.031. In our variable plot, the coordinates of rs4472504 were (-0.2113, -0.082), (-0.0987, 0.3589), (-0.028, 0.1049), and (-0.0995, -0.1101) in each group, respectively, whereas the coordinates of hypertension were (0.4645, 0.1513), (0.364, -0.364)

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0.1043), (0.4955, 0.0146), and (0.5504, -0.0219). Therefore, we cannot conclude that the relationship between a single SNP and a disease cannot be identical to the power of its influence on each axis.

Discussion

In this paper, we have reviewed how to analyze multi-block datasets—in particular, using genomewide data: the KARE dataset. We conducted GCCA in order to compare the relationship between and within multi-blocks. To see the relationship between variables, we used SAS [18] to do GCCA and R to visualize the relationships between and within multi-block data in four different subgroups: group 1, group 2, group 3, and group 4. In the relationship between blocks, we could reveal that there existed a stronger association between second canonical variables than between first canonical variables. In addition, we found that the relationship between SNP block and Phenotype block was the strongest, whereas the relationship between SNP block and Disease block was not remarkable. To see the relationship within variables, we made plots that showed how much each variable contributed to the canonical variable. Some SNP variables showed distinguishable influence among variable blocks, but most SNP variables did not show big difference. Phenotype variables, however, were distinguished by each group and showed dramatic differences between groups. In this paper, GCCA was applied to the preselected SNP set which relied on previous literatures, however, further research could be started from screening SNPs which are associated with phenotype or disease status.

We had limitations, in that different types of data, including both discrete and continuous data, can lead to unsuccessful results in GCCA. However, the analysis regarding SNP, phenotype, and disease at the same time would be meaningful itself and can even be more productive when we add pathway assumption in the association analysis. Therefore, further research of this topic should focus on robust generalized canonical correlation analysis, which could function regardless of datatype, and on the relationship that we would like to specify.

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Authors' contribution

Conceptualization: MP Data curation: IJ, WC, MP

Formal analysis: IJ, WC Funding acquisition: MP Methodology: IJ, WC, MP Writing – original draft: IJ, WC Writing – review & editing: IJ, MP

Conflicts of Interest

No potential conflicts of interest relevant to this article was reported.

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