Kernel canonical correlation analysis for assessing gene–gene interactions and application to ovarian cancer

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Although single-locus approaches have been widely applied to identify disease-associated single-nucleotide polymorphisms (SNPs), complex diseases are thought to be the product of multiple interactions between loci. This has led to the recent development of statistical methods for detecting statistical interactions between two loci. Canonical correlation analysis (CCA) has previously been proposed to detect gene–gene coassociation. However, this approach is limited to detecting linear relations and can only be applied when the number of observations exceeds the number of SNPs in a gene. This limitation is particularly important for next-generation sequencing, which could yield a large number of novel variants on a limited number of subjects. To overcome these limitations, we propose an approach to detect gene–gene interactions on the basis of a kernelized version of CCA (KCCA). Our simulation studies showed that KCCA controls the Type-I error, and is more powerful than leading gene-based approaches under a disease model with negligible marginal effects. To demonstrate the utility of our approach, we also applied KCCA to assess interactions between 200 genes in the NF-κB pathway in relation to ovarian cancer risk in 3869 cases and 3276 controls. We identified 13 significant gene pairs relevant to ovarian cancer risk (local false discovery rate < 0.05). Finally, we discuss the advantages of KCCA in gene–gene interaction analysis and its future role in genetic association studies.

Keywords: association studies; canonical correlation; gene–gene interaction; kernel methods

INTRODUCTION

Genome-wide association studies (GWAS) have identified hundreds of loci that harbor genetic variants that influence predisposition to a particular phenotype. Such studies involve the characterization of a large number of single-nucleotide polymorphisms (SNPs) across the genome and comparison of the frequency of variants across disease states. Initial GWAS analysis strategies involved single locus models, whereby individual markers were tested independently for association with a given phenotype. Although this approach has successfully identified regions of disease susceptibility, some contend it has failed to fully explain the heritability of complex phenotypes. As common complex diseases and traits are thought to be a result of complex interactions and multiple low-penetrance variants, multi-locus SNP models, as opposed to single SNP models, may better capture the true underlying genotypic–phenotypic relationship.

One strategy for multi-locus modeling is to jointly model the effects all SNPs within a given gene (eg, multivariable logistic regression models). However, this approach may lack power as the degrees of freedom of the model could be large and may require filtering or shrinkage approaches. Another drawback to the joint modeling of multiple SNPs within a gene is a possible model fitting issues due to multicollinearity between SNPs (ie, linkage disequilibrium (LD)), as well as the lack of inclusion of LD information in the analysis. Recently, this idea of gene-level analysis has led to the concept of gene–gene interaction analysis, as opposed to SNP–SNP interaction approaches. Gene–gene interactions are not only hypothesized to have a large role in explaining missing heritability, they can also serve to provide biological information through construction of novel gene pathway topologies. Although classically gene–gene interactions have been defined statistically as deviance from additive marginal effects, such as in the case of logistic regression model, this type of model is limiting with respect to statistical power. Moreover, the results of such SNP–SNP interaction analyses lack clear biological interpretability.

Zhao et al6 proposed testing for interactions between two unlinked loci using measures of LD, which can be extended to case-control design by comparing such measures across case or disease status. This concept was adapted to gene-level analysis by Peng et al7 which used a Wald-type U-statistic based on canonical correlation analysis8 (CCA) to detect gene–gene coassociation in case-control studies. As an LD-based procedure, the CCA approach obtains the maximal correlation of the linear combinations of the SNPs, coded as 0, 1 or 2 in terms of the minor allele, between two genes across case-control status, and tests whether the difference in the first canonical

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The transformation of the canonical correlation analyses such as SKAT12,13 and SPA-3G.14 Kernelized version of ovarian cancer to determine gene–gene interactions between genes gene interaction detection by a simulation study. Finally, we apply our statistical and computational issues that accompany this approach, CCA and its kernelized version. We further outline and address the case-control study of complex phenotypes. We briefly discuss sample spaces.

CCA (KCCA) provides a straightforward generalization of CCA to reduced kernel representations statistic based upon an analog of the Fisher variance stabilizing transformation (see Appendices I–III in Supplementary Material for details). We then define a kernel canonical correlations for cases and controls between genes both case and control status. For given genes control status, we use KCCA to generate measures of genetic coassociation for

\[ T_{lm} = \frac{\hat{x}_{lm}^{2} - \hat{x}_{lm}^{2}}{\text{var}(\hat{x}_{lm}^{2}) + \text{var}(\hat{x}_{lm}^{2})} \]

which is asymptotically distributed as \( N(0, 1) \) under the null hypothesis that \( \hat{x}_{lm}^{2} = \hat{x}_{lm}^{2} \), and the cases and controls are independent. For estimating the transform variances \( \text{var}(\hat{x}_{lm}^{2}) \) and \( \text{var}(\hat{x}_{lm}^{2}) \), we apply a robust resampling procedure, the trimmed jackknife16 (Appendix IV in Supplementary Materials).

**Multiple testing**

For applications involving exhaustive hypothesis testing of all pairwise gene comparisons in a given gene list, multiple testing and test statistic correlation become problematic issues. To address both of these directly, we apply Efron's empirical null method17 for estimating the local false discovery rate (LFDR) for each hypothesis test conducted.

**Simulation study**

To assess the properties of our KCCA procedure for gene–gene interaction testing, we consider simulation studies that evaluate type-I error control and power. We generated a population of haplotypes for two genes using real genotypic data from our case study. fastPHASE18 was applied to the genotypes from the controls to estimate haplotypes for two genes of comparable size (25.9 and 30.6 kb), followed by use of HapSim19 to simulate 10,000 haplotypes for each gene. The respective numbers of polymorphic sites for each gene were 79 and 92. Genotype data for a hypothetical individual were simulated by combining pairs of randomly selected haplotypes for each gene. Let \( m_{ij} \) represent the derived minor allele frequency (MAF) from our simulated haplotype populations for marker \( j \) in gene \( i \). Next, we randomly selected a fixed number of minor alleles (\( m_{ij} > 0.05 \)) markers to be causal for each gene. We then used a similar approach to effect size definition used by Wu et al.,20 in which effects are a function of the MAF. Let \( \beta_{ms}(m_{ij}, m_{ig}) \) be the interaction effect, such that \( \beta_{ms}(m_{ij}, m_{ig}) = \frac{\log(2m_{ij}m_{ig})}{\log_{10}(m_{ij}m_{ig})} \). Here \( t \) defines the maximum possible interaction effect. For example, for two markers with MAF = 0.20 and \( \tau = 5 \), the interaction effect is \( \beta_{ms}(m_{ij}, m_{ig}) = 0.141 \).

Given the difficulty in genome-wide detection of gene–gene interactions without the presence of marginal effects, we only considered disease models with solely epistatic effects. We defined the probability of being a case conditional on genotype \( P(Y=1 \mid X_{1}, X_{2}) \) via a logistic regression framework, such that

\[ \logit(P(Y=1 \mid X_{1}, X_{2})) = \beta_{0} + \sum_{j=1}^{m} \sum_{k=1}^{m} \beta_{ms}(m_{ij}, m_{ik}) x_{ij} x_{ik} \]

where \( \Omega_{i} \) and \( \Omega_{j} \) define the subsets of markers which are causal, and \( \beta_{0} \) represents the disease prevalence. Sampling of cases and controls was then completed from a sufficiently large number of simulated genotype-phenotype pairs.

To comparatively evaluate the performance of our KCCA method, we included additional methods on the basis of similar analysis principles: the original CCA-based approach, PC-Logistic regression22 (PC-LR), and a composite-LD method23 (CLD). PC-LR obtains principal components from SNP measurements for each gene, which are then fit in simple logistic regression. The CLD method is a covariance-based approach, which evaluates the difference between block interactions across case-control status. Additional details for each approach can be found in their respective publications. For PC-LR, we evaluate the significance of the interaction coefficient between the first principal component of each gene, and for the CLD approach we use 5000 permutations to characterize the reference distribution of the test statistic. All declarations of statistical significance are made at an \( \alpha \)-level of 0.05. For both Type-I error and power simulations, we consider whether or not explicit marginal effects are included in the disease model. Each simulation scenario is conducted with case-controls status sample sizes of 500, 1000, and 1500, with a total of 1000 iterations each.

**MATERIALS AND METHODS**

**Data definition**

Let \( s_{g} \) be the number of SNPs corresponding to the gth gene in a given gene list of size \( G \). Define \( x_{g}^{O} \) to be the genotype value for the jth SNP in gene \( g \) in the kth case subject, for \( g = 1, \ldots, G \), \( j = 1, \ldots, s_{g} \), and \( k = 1, \ldots, N_{D} \), where \( x_{g}^{O} \in \{0, 1, 2\} \) is the number of copies of the minor allele for SNP \( j \) in gene \( g \) under an assumed joint additive-additive genetic model. Similarly, we define \( x_{g}^{C} \) for the control subjects, where \( k = 1, \ldots, N_{C} \). This genotypic information can in turn be represented by the respective \( N_{D} \times s_{g} \) and \( N_{C} \times s_{g} \) matrices \( X_{g}^{O} \) and \( X_{g}^{C} \). These matrices may also contain adjusted genotype values that have been corrected for various covariates and population stratification.

**Hypothesis testing**

To test whether there is a statistical interaction between two genes across case-control status, we use KCCA to generate measures of genetic coassociation for both case and control status. For given genes \( l \neq m \in \{1, \ldots, G\} \), such that \( l \neq m \), consider the genotype matrices \( X_{l}^{O} \), \( X_{l}^{C} \), \( X_{m}^{O} \), \( X_{m}^{C} \) and \( X_{l,m}^{C} \) with corresponding reduced kernel representations \( K_{l}^{O}, K_{l}^{C}, K_{m}^{O}, K_{m}^{C}, K_{l,m}^{C} \) (see Appendices I–III in Supplementary Material for details). We then define a statistic based upon an analog of the Fisher variance stabilizing transformation of the Pearson’s correlation coefficient,15 given as

\[ z(r) = \frac{1}{2} \left\{ \log(1 + r) - \log(1 - r) \right\}. \]

The transformation of the canonical correlation \( z_{lm} \) is written as \( z_{lm}^{K} = z_{lm}^{C} \), which is approximately distributed as standard normal. A Wald-type statistic for assessing the statistical significance of the difference in gene–gene coassociation between cases and controls for genes \( l \) and \( m \) is defined by Peng et al2 as

\[ T_{lm} = \frac{\hat{x}_{lm}^{2} - \hat{x}_{lm}^{2}}{\text{var}(\hat{x}_{lm}^{2}) + \text{var}(\hat{x}_{lm}^{2})} \]

which is asymptotically distributed as \( N(0, 1) \) under the null hypothesis that \( \hat{x}_{lm}^{2} = \hat{x}_{lm}^{2} \), and the cases and controls are independent. For estimating the transform variances \( \text{var}(\hat{x}_{lm}^{2}) \) and \( \text{var}(\hat{x}_{lm}^{2}) \), we apply a robust resampling procedure, the trimmed jackknife16 (Appendix IV in Supplementary Materials).
Ovarian cancer study
We applied the KCCA approach to detect gene–gene interaction within the NF-κB pathway, using data from a case-control study of invasive epithelial ovarian cancer as part of the GAME-ON Follow-up Ovarian Cancer Genetic Association and Interaction Studies collaboration (described elsewhere). Participants were enrolled in the Mayo Clinic Ovarian Cancer Study, North Carolina Ovarian Cancer Study, the Tampa Bay Ovarian Cancer Study, the Toronto Ovarian Cancer Study, the National Cancer Institute Ovarian Case-Control Study in Poland, the UK Ovarian Cancer Population Study, the Studies of Epidemiology and Risk Factors in Cancer Heredity Ovarian Cancer Study, the Familial Ovarian Cancer Registry Study, and the Royal Marsden Hospital Ovarian Cancer Study. Study protocols were approved by the appropriate institutional review board or ethics panel, and all patients provided written informed consent. Genotypes were from Illumina (San Diego, CA, USA) 610-Quad SNP arrays, with imputation to HapMap v26 using MACH.26 For 200 autosomal genes within the NF-κB pathway, this resulted in ~13,000 observed or imputed markers available on 3869 cases and 3276 controls of European descent. For genotyped markers, we coded genotypes as 0, 1, or 2 in terms of the number of observed minor alleles; for imputed markers, we used the expected genotype or ‘dosage’. Marker assignment to genes was determined on the basis of NCBI-build 36 gene location data, using a 20-kb buffer region on both the 5’ and 3’ ends of the defined gene location. Information on location, size, and number of SNPs in each of the 200 autosomal genes can be found in Supplementary Table S1.

To address the effects of possible confounding variables, we adjusted the genotypes for age, study site, and the first five principal components from an eigen analysis. Each unique gene pair was tested using the KCCA procedure, resulting in \( \binom{200}{2} = 19900 \) total hypothesis tests. For purposes of comparison, we also applied the CCA-based procedure defined by Peng et al.27 To the data, we used 1000 bootstraps for variance estimation. Comparisons involving gene pairs with overlapping regions were removed from our analysis to avoid complications involving shared marker data.

RESULTS
Type-I error
For our simulations, we considered six levels of trimming within the jackknife procedure for SE estimation to determine which level was appropriate. A plot of the empirical Type-I error rate for all trim levels across each sample size are found in Figure 1. The KCCA results derived from \( \omega = 0.15 \) yield near nominal Type-I error rate levels across all sample sizes. Detailed statistics on the empirical distributions of the test statistics can be found in Table 1. These results indicate that the test statistic follows the assumed standard normal distribution under the null.

Power
For our power simulations, we set \( \tau = 5 \) and randomly selected five markers from each gene to be causal at each iteration, performing all KCCA tests with \( \omega = 0.15 \). Bar graphs of the results are found in Figure 2. From this plot, we observe that KCCA outperforms the other methods across all sample sizes, particularly the CCA approach. As the poor performance of CCA is likely due to issues with overfitting, we considered additional simulations where sample sizes were fixed at 1000 and the number of markers per gene was set to values of 10, 20, 30, 40, or 50, randomly subsetted from the total number of markers used in our simulations. Keeping all the other settings of our simulation design the same, the empirical power for each method across number of markers per gene is found in Table 2. Although the resulting power is much closer for low marker numbers, the KCCA approach is still consistently more powerful than the remaining methods.

Application to ovarian cancer study
For the ovarian cancer study of the NF-κB pathway, we identified 13 statistically significant (IFDR < 0.05) gene pairs of interest (Table 3) in applying our KCCA method, using the trimmed jackknife SE estimate with \( \omega = 0.15 \), with the top coassociated gene pair with case-control status occurring between CASP8 and MAP3K3. Application of the CCA-based procedure resulted in 37 significant gene pairs; however, none of these overlapped with those detected by the KCCA method.

To explore one of the top coassociation hits at the SNP level, we analyzed the CASP8–MAP3K3 interaction by generating pairwise marker Pearson’s correlations. These are presented by case-control status in Figure 3 as gradient colorized images of the correlation coefficients. The figure demonstrates that although the Pearson’s correlation coefficients themselves are small (|\( r \)| < 0.10), there exist distinct differences between the two correlation structures. Of note is the discrepancy across case-control status with the correlation between marker rs12940055 in MAP3K3 (located at bp 59075874) and a large number of SNPs toward the 5’ end of CASP8, indicated by the horizontal band in the lower half of the correlation plots.

DISCUSSION
The identification of interactions between genes and their impact on complex diseases is adding to our understanding the genetic component. Although SNP-based interaction analysis methods are relatively well developed, the large number of SNPs in association studies makes exhaustive pairwise SNP-SNP analyses increasingly infeasible. By addressing the problem at the gene level, the scope of the analyses is not only computationally tenable, but also reduced to a biologically interpretable unit of interest. Peng et al. presented a novel statistical method that allows for such analyses in their CCA statistic. However, as the number of genotyped markers increases, the application of CCA may be inappropriate, because the number of genotyped SNPs may exceed the number of observations, especially for smaller experimental designs. This is of particular concern with post-GWAS whole exome and genome sequencing association studies, which will afford the characterization of additional variants unmeasured by current genotyping platforms.

In this article, we have presented a KCCA for gene–gene interaction analysis that not only addresses concerns of dimensionality, but also allows the flexibility to detect nonlinear correlations between genes. We have demonstrated that, using the appropriate SE estimation procedure, the KCCA test statistic exhibits near nominal levels of type-I error rate control and competitive power performance in our
Table 1  KCCA null simulation results for various trimming values $\omega$, which includes the $P$-value for the Kolmogorov-Smirnov test for normality (KS), the empirical SD and mean of the simulated test statistic distribution, as well as the realized Type-I error rate rejecting at an $\alpha$-level of 0.05

| $\omega$ | KS | SD | Mean | Type I | KS | SD | Mean | Type I | KS | SD | Mean | Type I |
|----------|----|----|------|--------|----|----|------|--------|----|----|------|--------|
| 0.00     | <0.001 | 0.633 | 0.002 | 0.001 | <0.001 | 0.643 | 0.001 | 0.003 | <0.001 | 0.757 | 0.002 | 0.006 |
| 0.05     | <0.001 | 0.798 | -0.001 | 0.012 | <0.001 | 0.801 | -0.001 | 0.016 | <0.001 | 0.938 | -0.001 | 0.021 |
| 0.10     | <0.001 | 0.904 | -0.003 | 0.033 | <0.001 | 0.902 | -0.002 | 0.027 | 0.010 | 1.038 | 0.002 | 0.036 |
| 0.15     | 0.281 | 1.007 | -0.003 | 0.540 | 0.810 | 1.008 | -0.004 | 0.046 | 0.538 | 1.144 | 0.004 | 0.054 |
| 0.20     | 0.002 | 1.107 | -0.001 | 0.081 | 0.002 | 1.114 | -0.005 | 0.072 | 0.001 | 1.250 | 0.005 | 0.078 |
| 0.25     | <0.001 | 1.201 | -0.001 | 0.107 | <0.001 | 1.220 | -0.005 | 0.095 | <0.001 | 1.363 | 0.004 | 0.110 |

Figure 2  Barplots of the empirical power at $\alpha$-level of 0.05 for the KCCA, CCA, PC-LR, and CLD gene–gene interaction methods, for sample sizes of 500, 1000, and 1500.

Table 2  Simulation result for empirical power, using $\alpha$ = 0.05 level significance testing, for the KCCA, CCA, PC-LR, and CLD testing procedures across varying numbers of markers per gene

| Total markers per gene | 10 | 20 | 30 | 40 | 50 |
|------------------------|----|----|----|----|----|
| KCCA                   | 0.852 | 0.818 | 0.799 | 0.779 | 0.757 |
| CCA                    | 0.428 | 0.097 | 0.012 | 0.002 | 0.000 |
| PC-LR                  | 0.570 | 0.575 | 0.552 | 0.545 | 0.544 |
| CLD                    | 0.724 | 0.570 | 0.371 | 0.305 | 0.234 |

Comparability with CCA statistic
Contrary to the findings of Peng et al., we have found that hypothesis testing using their CCA statistic with bootstrap variance estimation can be quite conservative, regardless of sample or feature size used, resulting in possible issues with reduced statistical power. This is evidenced in our power simulations where CCA power results for certain simulation conditions result in power below even nominal Type-I error rates. Upon investigation of this discrepancy, we found that the bootstrap variance estimates were quite large in comparison with their KCCA counterparts.

An additional benefit of our procedure is that the assumptions of CCA include multivariate normality of the observations, which is clearly violated by the discrete nature of genotype calls if no adjustments are made. Our method, however, involves low dimensional projections of kernelized observations, which has been shown to be approximately Gaussian. Thus, our KCCA procedure is also more consistent with the distributional assumptions of CCA.

Ovarian cancer findings
The analysis of the FOCI data using the KCCA procedure yielded biologically interesting results, with many of the top gene pairs sharing some functional basis. CASP8 and MAP3K3 are integral members of the tumor necrosis factor pathway, and IL1A and IL1B both code for proinflammatory cytokines and have been jointly associated with lung cancer. With 13 statistically significant findings, there are also several novel interactions that may warrant further investigation.
We argue the lack of congruency between the results of the CCA-based procedure and our kernelized version is due, in large part, to the lack of any dimensional reduction in the CCA. Although the sample sizes used in our ovarian application analyses are large enough to satisfy most conventional notions of appropriate observation-to-feature ratios for accurate canonical correlation estimation, our simulations indicate that the CCA method suffers greatly from overfitting when there are a relatively large number of genotyped markers in given genes.

One significant gene pair of concern in the NF-κB–KCCA interaction analysis is \( \text{IL1A-IL1B} \), which includes genes that are positionally adjacent to one another. Taking into account their respective buffer regions, only 4.4 kb separates the two genes. The significance of this gene pair could be evidence of instability of the method for gene pairs that are in high LD with each other because of locational proximity. However, gene pairs of this nature are also often closely linked functionally. Moreover, there are counter examples in our analysis of neighboring gene pairs that are statistically not significant (e.g., \( \text{LTBR-TNFRSF1A} \)). Regardless, we recommend caution in the interpretation of such results.

### Table 3 Detailed results of the significant KCCA gene–gene coassociations for analysis of ovarian cancer risk of significant (lFDR ≤ 0.05) gene–gene interactions from FOCI analysis ranked by estimated lFDR

| Gene 1   | Gene 2   | \( z_{12}^2 \) | \( z_{21}^2 \) | \( T_{12} \) | \( P\text{-value} \) | lFDR  | T12  | \( P\text{-value} \) |
|----------|----------|----------------|----------------|-------------|-----------------|-------|------|------------------|
| CASP8    | MAP3K3   | 0.0875         | 0.0074         | 14.2897     | <1.00E-16       | 3.38E-12 |      | 0.7248           | 0.46853         |
| IL1A     | IL1B     | 1.7243         | 1.9167         | -12.8110    | <1.00E-16       | 1.58E-15 |      | -0.7284          | 0.92525         |
| MAP3K3   | TAB1     | 0.0976         | 0.0196         | 8.1708      | 2.22E-16        | 8.53E-06 |      | -0.1093          | 0.82548         |
| MAPK8    | FEL13    | 0.1146         | 0.0462         | 8.2271      | 2.22E-16        | 7.11E-06 |      | -0.0947          | 0.92449         |
| AZ12     | IKBK8    | 0.1049         | 0.0344         | 9.2402      | <1.00E-16       | 5.65E-08 |      | 0.2640           | 0.79171         |
| 7BKBP1   | TLR10    | 0.0436         | 0.0989         | -6.8641     | 6.69E-12        | 0.008161 |      | -0.1386          | 0.88972         |
| HEXIM1   | MYD88    | 0.1073         | 0.0524         | 6.8256      | 8.75E-12        | 0.001646 |      | -0.3888          | 0.69736         |
| PIK3CB   | TAB1     | 0.1025         | 0.0317         | 10.6548     | <1.00E-16       | 2.59E-11 |      | -0.2520          | 0.80098         |
| BIRC3    | MAP3K3   | 0.0763         | 0.0194         | 5.7655      | 8.14E-09        | 0.028977 |      | 0.7772           | 0.43702         |
| MAP3K1   | TRAF7    | 0.1125         | 0.0465         | 5.5000      | 3.80E-08        | 0.049916 |      | -1.0316          | 0.30223         |
| BIRC3    | TNFRSF13C| 0.0601         | 0.0001         | 10.8523     | <1.00E-16       | 8.74E-12 |      | -0.1986          | 0.84251         |
| MAP3K1   | MAP3K8   | 0.1032         | 0.0377         | 6.1801      | 6.41E-10        | 0.010925 |      | 0.3529           | 0.72412         |
| PYCARD5  | TAF4     | 0.0231         | 0.1012         | -6.5902     | 4.39E-11        | 0.016345 |      | 0.0313           | 0.97502         |

**Abbreviations:** CCA, canonical correlation analysis; FOCI, Follow-up Ovarian Cancer Genetic Association and Interaction Studies; KCCA, kernelized version of CCA; lFDR, local false discovery rate. Includes the Fisher-transformed maximal kernel canonical correlation values for cases and controls, the resulting test statistic, \( P\text{-values} \), and lFDR estimates for KCCA, as well as results for the CCA analysis.

**Figure 3** Colorized image plot of Pearson’s correlation values between SNPs for CASP8-MAP3K3 coassociation for cases (left) and controls (right). The axes depict the genomic position of the markers on the respective genes.

**Figure 4** Barplots of the empirical power at \( \alpha \)-level of 0.05 for the KCCA, CCA, PC-LR, and CLD gene–gene interaction methods, under sample sizes of 500, 1000, and 1500 with the inclusion of statistically significant marginal effects.

We argue the lack of congruency between the results of the CCA-based procedure and our kernelized version is due, in large part, to the lack of any dimensional reduction in the CCA. Although the sample sizes used in our ovarian application analyses are large enough to satisfy most conventional notions of appropriate observation-to-feature ratios for accurate canonical correlation estimation, our simulations indicate that the CCA method suffers greatly from overfitting when there are a relatively large number of genotyped markers in given genes.

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Conclusions and future development

Our KCCA algorithm simultaneously supplies dimensionality reduction and nonlinear coassociation analysis for high-dimensional SNP data, providing a powerful framework for detecting statistical epistasis at the gene level. Moreover, this type of analysis can isolate gene pairs of interest for follow-up analysis without being burdened by the multiple testing corrections necessary for genome-wide SNP–SNP pairwise interaction analysis. This is particularly relevant for next-generation sequencing applications, which may interrogate all possible SNPs through whole genome sequencing.

Although we have argued that the KCCA procedure for detecting gene–gene interaction possesses many advantages over the previously proposed CCA statistic, there is also room for improvement and generalizability of our approach. The use of the Gaussian kernel function is a robust selection; however, other kernel functions may be more appropriate for the specific data type, particularly if there are no adjustments for covariate data.34 The procedure itself may also be modified in a variety of ways, including the use of sparse canonical correlation22,23 and multigene interaction analysis with generalized canonical correlation,24 and further exploring the resampling procedures used in the SE estimation. Finally, a less computationally intensive alternative to KCCA may be a kernelized variant of principal correlation,25 which could be considered for more demanding analyses such as genome-wide interrogation.

CONFLICT OF INTEREST

The authors declare no conflict of interest.

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