# GeneGeneInteR vignette Statistical analysis of the interaction between a pair of genes. 

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This vignette presents the technical details of the statistical procedure implemented in the package. Readers that would like to have a global overview of the main functions and tools proposed in the package are encouraged to read the vignette VignetteGeneGeneInteR_Introduction.

## 1 Introduction

In this vignette we consider statistical procedures to test for the interaction between two genes in susceptibility with a binary phenotype, typically a case/control disease status. Let $Y \in\{0,1\}$ be the phenotype, where $Y=0$ stands for a control and $Y=1$ a case, and $X_{1}$ and $X_{2}$ be the two genes for which the interaction is tested.

Let consider a sample of $n$ individuals with $n_{c}$ controls and $n_{d}$ cases $\left(n_{c}+n_{d}=n\right)$ and $\mathbf{Y}=\left[y_{1}, \ldots, y_{n}\right]^{\prime}$ the vector of the observed binary phenotypes. Each gene is a collection of respectively $m_{1}$ and $m_{2}$ SNPs. The observed genotypes for gene $X_{1}$ can be represented by a $n \times m_{1}$ matrix: $\mathbf{X}_{1}=\left[x_{i j}^{1}\right]_{i \in 1 \ldots n ; j \in 1 \ldots m_{1}}$ where $x_{i j}^{1} \in\{0 ; 1 ; 2\}$ is the number of copies of the minor allele for SNP $j$ carried by individual $i$. A similar representation is used for gene $X_{2}$ where $\mathbf{X}_{\mathbf{2}}$ is a $n \times m_{2}$ matrix. Let us further introduce $\mathbf{X}_{\mathbf{1}}^{\mathbf{c}}$ and $\mathbf{X}_{\mathbf{2}}^{\mathbf{c}}$ the matrices of observed genotypes among controls for gene 1 and 2 and $\mathbf{X}_{\mathbf{1}}^{\mathbf{d}}$ and $\mathbf{X}_{\mathbf{2}}^{\mathbf{d}}$ among cases for both genes. Thus $\mathbf{X}_{1}^{\mathbf{c}}$ is a $n_{c} \times m_{1}$ matrix, $\mathbf{X}_{1}^{\mathbf{d}}$ a $n_{d} \times m_{1}$ matrix, $\mathbf{X}_{2}^{\mathbf{c}}$ a $n_{c} \times m_{2}$ matrix and $\mathbf{X}_{2}^{\mathbf{d}}$ a $n_{d} \times m_{2}$ matrix. A general setup of the observed values can be presented as follows:

In our package we proposed 10 methods for testing interaction at the gene level. These 10 methods are all based on three main parameters: Y, a numeric or factor vector with exactly two distinct values, G1 and G2 two SnpMatrix objects as proposed by the R Bioconductor package snpStats. Our implementation is illustrated by the dataset gene. pair provided with the GeneGeneInteR package and summarized in the following command lines:

```
> library("GeneGeneInteR")
> data("gene.pair")
> head(gene.pair$Y)
```

[1] HealthControl HealthControl HealthControl HealthControl HealthControl Levels: HealthControl RheumatoidArthritis

```
> gene.pair$G1
```

```
A SnpMatrix with }453\mathrm{ rows and 8 columns
Row names: Id1 ... Id453
Col names: rs1491710 ... rs2298849
> gene.pair$G2
A SnpMatrix with 453 rows and 4 columns
Row names: Id1 ... Id453
Col names: rs2057094 ... rs1005753
```

The 10 methods implemented in our package can be divided into two main families: 6 methods based on a multidimensional modeling of the interaction at the gene level and 4 methods that combine interaction tests at the SNP level into a single test at the gene level.

## 2 Multidimensional methods at the gene level

In the GeneGeneInteR package, 6 multidimensional methods have been implemented that are based on:

- Principal Components Analysis - PCA.test function,
- Canonical Correlation Analysis - CCA.test function,
- Kernel Canonical Correlation Analysis - KCCA.test function,
- Composite Linkage Disequilibrium - CLD.test function,
- Partial Least Square Path Modeling - PLSPM.test function,
- Gene-Based Information Gain Method - GBIGM.test function.

In the remainder of this section, technical and practical details are given regarding these 6 methods.

### 2.1 PCA-based

In the PCA-based method, a likelihood ratio test is performed to compare the model $\mathcal{M}_{\text {Inter }}$ to the model $\mathcal{M}_{\text {No }}$, where $\mathcal{M}_{\text {Inter }}$ is defined by:

$$
\operatorname{logit}\left(\mathbb{P}\left[Y=1 \mid P C_{X_{1}}^{1} \ldots P C_{X_{1}}^{n_{1}}, P C_{X_{2}}^{1} \ldots P C_{X_{2}}^{n_{2}}\right]\right)=\beta_{0}+\sum_{i=1}^{n_{1}} P C_{X_{1}}^{i}+\sum_{j=1}^{n_{2}} P C_{X_{2}}^{j}+\sum_{i=1}^{n_{1}} \sum_{i=2}^{n_{2}} P C_{X_{1}}^{i} P C_{X_{2}}^{j}
$$

and $\mathcal{M}_{N o}$ by:

$$
\operatorname{logit}\left(\mathbb{P}\left[Y=1 \mid P C_{X_{1}}^{1} \ldots P C_{X_{1}}^{n_{1}}, P C_{X_{2}}^{1} \ldots P C_{X_{2}}^{n_{2}}\right]\right)=\beta_{0}+\sum_{i=1}^{n_{1}} P C_{X_{1}}^{i}+\sum_{j=1}^{n_{2}} P C_{X_{2}}^{j}
$$

In models $\mathcal{M}_{\text {Inter }}$ and $\mathcal{M}_{N o}, P C_{X_{1}}^{i}$ and $P C_{X_{2}}^{j}$ are the $i^{\text {th }}$ principal component of $\mathbf{X}_{\mathbf{1}}$ and the $j^{\text {th }}$ principal component of $\mathbf{X}_{\mathbf{2}}$. The number of principal components, $n_{1}$ and $n_{2}$, kept in the interaction test is determined by the percentage of inertia retrieved by the PCA. Such a percentage is defined by the user and corresponds to the threshold parameter.

In our package, two distinct Principal Component decomposition are provided by the functions PCA.test via the argument method. With method="Std", dataset is standardized using variables' standard deviation while with method="GenFreq", dataset is standardized using standard deviation under Hardy-Weinberg equilibrium, as proposed in the snpStats package.

When the percentage of inertia asked by the user is high, the number of PCs can be important and fitting logistic models $\mathcal{M}_{\text {Inter }}$ and $\mathcal{M}_{N o}$ is likely to fail. In that case, the number of PCs in each gene is iteratively reduced until convergence of the glm function for fitting models $\mathcal{M}_{\text {Inter }}$ and $\mathcal{M}_{N o}$.

The following lines provide an example of the PCA.test function:

```
> PCA.test(Y=gene.pair$Y, G1=gene.pair$G1, G2=gene.pair$G2,threshold=0.7,
+ method="GenFreq")
```

Gene-based interaction based on Principal Component Analysis - GenFreq

```
data: gene.pair$Y and (gene.pair$G1 , gene.pair$G2)
```

Deviance $=8.2157, \mathrm{df}=6.0$, threshold $=0.7, \mathrm{p}$-value $=0.2227$
alternative hypothesis: true deviance is greater than 0
sample estimates:
Deviance without interaction Deviance with interaction
615.2977607 .0821
> PCA.test(Y=gene.pair\$Y, G1=gene.pair\$G1, G2=gene.pair\$G2,threshold=0.7,

+ method="Std")
Gene-based interaction based on Principal Component Analysis - Std
data: gene.pair\$Y and (gene.pair\$G1, gene.pair\$G2)
Deviance $=8.5074, \mathrm{df}=6.0$, threshold $=0.7, \mathrm{p}$-value $=0.2032$
alternative hypothesis: true deviance is greater than 0
sample estimates:
Deviance without interaction Deviance with interaction
615.0911
606.5837


### 2.2 Canonical Correlation Analysis (CCA)

The CCA test is based on a Wald-type statistic defined as follows (see Peng et al., 2010 for details):

$$
U_{C C A}=\frac{z_{d}-z_{c}}{\sqrt{\mathbb{V}\left(z_{d}\right)+\mathbb{V}\left(z_{c}\right)}}
$$

where $z_{d}=\frac{1}{2}\left(\log \left(1+r_{d}\right)-\log \left(1-r_{d}\right)\right)$ and $z_{c}=\frac{1}{2}\left(\log \left(1+r_{c}\right)-\log \left(1-r_{c}\right)\right)$ with $r_{d}$ the maximum canonical correlation coefficient between $\mathbf{X}_{1}^{\mathbf{d}}$ and $\mathbf{X}_{\mathbf{2}}^{\mathbf{d}}$ and $r_{c}$ the maximum canonical correlation coefficient between $\mathbf{X}_{\mathbf{1}}^{\mathbf{c}}$ and $\mathbf{X}_{2}^{\mathbf{c}}$ computed for controls $(Y=0)$. As suggested by Peng et al., 2010, the sampled variances $\mathbb{V}\left(z_{d}\right)$ and $\mathbb{V}\left(z_{c}\right)$ were evaluated by applying a bootstrapping method. The number of bootstrap sample used to estimate $\mathbb{V}\left(z_{d}\right)$ and $\mathbb{V}\left(z_{c}\right)$ is determined by the n . boot argument. P-value is then obtained by noting that under the null hypothesis $U_{C C A} \sim \mathcal{N}(0,1)$.

CCA based gene-gene interaction is implemented in the CCA.test function and mainly depends on the cancor function from the Stats package R Core Team, 2016.

```
    R> set.seed (1234)
> CCA.test(Y=gene.pair$Y, G1=gene.pair$G1, G2=gene.pair$G2,n.boot=500)
    Gene-based interaction based on Canonical Correspondance Analysis
data: gene.pair$Y and (gene.pair$G1 , gene.pair$G2)
CCU = 0.60304, n.boot = 500, p-value = 0.5465
alternative hypothesis: true CCU is not equal to 0
sample estimates:
    z0 z1
0.2940799 0.2414700
```


### 2.3 Kernel Canonical Correlation Analysis (KCCA)

The KCCA based test provides a generalization of CCA test to detect non-linear co-association between $\mathbf{X}_{\mathbf{1}}$ and $\mathbf{X}_{\mathbf{2}}$ Yuan et al., 2012, Larson and Schaid, 2013 and is based on the following Wald-type statistic:

$$
U_{K C C A}=\frac{k z_{d}-k z_{c}}{\sqrt{\mathbb{V}\left(k z_{d}\right)+\mathbb{V}\left(k z_{c}\right)}}
$$

where $k z_{d}=\frac{1}{2}\left(\log \left(1+k r_{d}\right)-\log \left(1-k r_{d}\right)\right)$ and $k z_{c}=\frac{1}{2}\left(\log \left(1+k r_{c}\right)-\log \left(1-k r_{c}\right)\right)$ with $k r_{d}$ the maximum kernel canonical correlation coefficient between $\mathbf{X}_{1}^{\mathbf{d}}$ and $\mathbf{X}_{2}^{\mathbf{d}}$ and $k r_{c}$ the maximum kernel canonical correlation coefficient between $\mathbf{X}_{1}^{\mathbf{c}}$ and $\mathbf{X}_{\mathbf{2}}^{\mathbf{c}}$.

Similar to the CCA test, $\mathbb{V}\left(k z_{d}\right)$ and $\mathbb{V}\left(k z_{c}\right)$ are estimated using bootstrap techniques Yuan et al., 2012, Larson and Schaid, 2013 and the p-value is obtained using the standard gaussian distribution of $U_{K C C A}$ under the null hypothesis. Since the performance of kernel methods strongly relates to the choice of kernel functions, the default is the Radial Basis kernel Function (RBF) owing to its flexibility in parameter specification. However, other kernel functions, such as linear, polynomial or spline kernels, can be used. Thus, in addition to the three arguments Y, G1 and G2, our implementation of the KCCA test proposes two optional arguments: n. boot that determines the number of bootstrap samples and kernel that provides the kernel function to be used. This kernel parameter is character string matching one of the kernel name provided by the kernlab package Karatzoglou et al., 2004 such as "rbfdot", "polydot", "tanhdot", "vanilladot", "laplacedot", "besseldot", "anovadot", "splinedot". Specific arguments, sigma, degree, scale, offsetand order, can also be passed to the kcca.test function in order to parameterized the kernel used in the analysis.

KCCA based gene-gene interaction test is implemented in the KCCA.test function and mainly depends on the kcca function from the kernlab package Karatzoglou et al., 2004.

```
> set.seed(1234)
> KCCA.test(Y=gene.pair$Y, G1=gene.pair$G1,G2=gene.pair$G2,
+ kernel="rbfdot",sigma = 0.05,n.boot=500)
    Gene-based interaction based on Kernel Canonical Correspondance
    Analysis
data: gene.pair$Y and (gene.pair$G1 , gene.pair$G2)
KCCU = 1.4055, n.boot = 500, p-value = 0.1599
alternative hypothesis: true KCCU is not equal to 0
sample estimates:
    z0 z1
    3.717346 -3.759154
> set.seed(1234)
> KCCA.test(Y=gene.pair$Y, G1=gene.pair$G1,G2=gene.pair$G2,
+ kernel="polydot",degree = 1, scale = 1, offset = 1)
    Gene-based interaction based on Kernel Canonical Correspondance
    Analysis
data: gene.pair$Y and (gene.pair$G1 , gene.pair$G2)
KCCU = 1.4106, n.boot = 100, p-value = 0.1584
alternative hypothesis: true KCCU is not equal to 0
sample estimates:
    z0 z1
4.161048-4.251702
```


### 2.4 Partial Least Square Path Modeling (PLSPM)

The PLSPM testing has been introduced by Zhang et al., 2013 and is based on the Wald-like statistic:

$$
U_{P L S P M}=\frac{\beta_{d}-\beta_{c}}{\sqrt{\mathbb{V}\left(\beta_{d}-\beta_{c}\right)}}
$$

where $\beta_{d}$ (resp. $\beta_{c}$ ) is the path coefficient between $\mathbf{X}_{\mathbf{1}}^{\mathbf{d}}$ and $\mathbf{X}_{\mathbf{2}}^{\mathbf{d}}$ (resp. $\mathbf{X}_{\mathbf{1}}^{\mathbf{c}}$ and $\mathbf{X}_{\mathbf{2}}^{\mathbf{c}}$ ). As quoted by Zhang et al., 2013], the distribution of $U_{P L S P M}$ is unknown and significance can be tested with bootstrapping method.

PLSPM based gene-gene interaction test is implemented in the PLSPM.test function and mainly depends on the plspm function from the plspm package Sanchez et al., 2015.

```
> set.seed(1234)
> PLSPM.test(Y=gene.pair$Y, G1=gene.pair$G1,G2=gene.pair$G2,n.perm=1000)
    Gene-based interaction based on Partial Least Squares Path Modeling
data: gene.pair$Y and (gene.pair$G1 , gene.pair$G2)
U = 4.0938, n.perm = 1000, p-value = 0.18
alternative hypothesis: true U is not equal to 0
sample estimates:
    beta0 beta1
-0.2125869 0.2434624
```


### 2.5 Composite Linkage Disequilibrium (CLD)

The CLD method, proposed in Rajapakse et al., 2012 is based on the normalized quadratic distance (NQD) and is defined as

$$
\delta^{2}=\operatorname{tr} \cdot\left((\tilde{D}-\tilde{C}) W^{-1}(\tilde{D}-\tilde{C}) W^{-1}\right)
$$

where $\tilde{D}, \tilde{C}$ and $W$ are three $\left(m_{1}+m_{2}\right) \times\left(m_{1}+m_{2}\right)$ matrices of the covariance between the whole set of SNPs that combines SNPs from both genes. More precisely, $\tilde{D}$ and $\tilde{C}$ are defined as follows:

$$
\tilde{D}=\left[\begin{array}{ll}
W_{11} & D_{12} \\
D_{21} & W_{22}
\end{array}\right] \quad \tilde{C}=\left[\begin{array}{ll}
W_{11} & C_{12} \\
C_{21} & W_{22}
\end{array}\right]
$$

where $W_{11}$ (resp. $W_{22}$ ) is the pooled estimate of the covariance matrix for $\mathbf{X}_{\mathbf{1}}$ (resp. $\mathbf{X}_{\mathbf{2}}, D_{12}\left(=D_{21}^{\prime}\right)$ and $C_{12}\left(=C_{21}^{\prime}\right)$ are the sample covariance matrix between the two genes estimated from $\left(\mathbf{X}_{\mathbf{1}}^{\mathbf{d}}, \mathbf{X}_{\mathbf{2}}^{\mathbf{d}}\right)$ and $\left(\mathbf{X}_{\mathbf{1}}^{\mathbf{c}}, \mathbf{X}_{\mathbf{2}}^{\mathbf{c}}\right)$ respectively. In more details, the sample covariance matrices in cases, denoted by $D$, and in controls, denoted by $C$, can be partitioned in 4 blocks as follows:

$$
D=\operatorname{Cov}\left(\mathbf{X}_{\mathbf{1}}^{\mathbf{d}}, \mathbf{X}_{\mathbf{2}}^{\mathbf{d}}\right)=\left[\begin{array}{ll}
D_{11} & D_{12} \\
D_{21} & D_{22}
\end{array}\right] \quad C=\operatorname{Cov}\left(\mathbf{X}_{\mathbf{1}}^{\mathbf{c}}, \mathbf{X}_{\mathbf{2}}^{\mathbf{c}}\right)=\left[\begin{array}{ll}
C_{11} & C_{12} \\
C_{21} & C_{22}
\end{array}\right]
$$

The pooled estimate of the covariance matrix, $W$, can thus been obtained by:

$$
W=\frac{n_{c} C+n_{d} D}{n_{c}+n_{d}}=\left[\begin{array}{ll}
W_{11} & W_{12} \\
W_{21} & W_{22}
\end{array}\right]
$$

Since the distribution of $\delta^{2}$ is not known under the null hypothesis, significance testing is performed using permutation tests, as proposed by Rajapakse et al., 2012. Such a test has been implemented in our package in the CLD.test function where the number of permutations is determined by the argument $n$.perm.

```
> set.seed(1234)
> CLD.test(Y=gene.pair$Y, G1=gene.pair$G1,G2=gene.pair$G2,n.perm=2000)
```

Gene-based interaction based on Composite Linkage Disequilibrium

```
data: gene.pair$Y and (gene.pair$G1 , gene.pair$G2)
CLD = 0.49257, n.perm = 2000, p-value = 0.8865
alternative hypothesis: true CLD is not equal to 0
sample estimates:
    CLD
0.4925654
```


### 2.6 Gene-Based Information Gain Method (GBIGM)

Introduced by Li et al., 2015, the GBIGM method is based on the information gain rate $\Delta R_{1,2} . \Delta R_{1,2}$ is defined as follows:

$$
\Delta R_{1,2}=\frac{\min \left(H_{1}, H_{2}\right)-H_{1,2}}{\min \left(H_{1}, H_{2}\right)}
$$

where $H_{1}, H_{2}, H_{1,2}$ are the conditional entropies, given the $\mathbf{Y}$, of $\mathbf{X}_{\mathbf{1}}, \mathbf{X}_{\mathbf{2}}$ and the pooled SNP set $\left(\mathbf{X}_{\mathbf{1}}, \mathbf{X}_{\mathbf{2}}\right)$ respectively. Assuming that $H($.$) is the classical entropy function, we have:$

$$
\begin{aligned}
H_{1} & =H\left(\mathbf{Y}, \mathbf{X}_{\mathbf{1}}\right)-H\left(\mathbf{X}_{\mathbf{1}}\right) \\
H_{2} & =H\left(\mathbf{Y}, \mathbf{X}_{\mathbf{2}}\right)-H\left(\mathbf{X}_{\mathbf{2}}\right) \\
H_{1,2} & =H\left(\mathbf{Y}, \mathbf{X}_{\mathbf{1}}, \mathbf{X}_{\mathbf{2}}\right)-H\left(\mathbf{X}_{\mathbf{1}}, \mathbf{X}_{\mathbf{2}}\right)
\end{aligned}
$$

Since the distribution of $\Delta R_{1,2}$ is unknown, the significance testing is performed by permutations as suggested by Li et al., 2015. The GBIGM method has been implemented in the GBIGM. test function and the number of permutations is defined by the argument n.perm.

```
> set.seed(1234)
> GBIGM.test(Y=gene.pair$Y, G1=gene.pair$G1,G2=gene.pair$G2,n.perm=2000)
    Gene-based interaction based on Gene-based Information Gain Method
data: gene.pair$Y and (gene.pair$G1 , gene.pair$G2)
DeltaR1,2 = 0.46441, n.perm = 2000, p-value = 0.441
alternative hypothesis: two.sided
sample estimates:
DeltaR1,2
0.4644093
```


## 3 From SNP-SNP interaction to Gene-Gene interaction testing

This section provides details of the four statistical methods that proposes a gene-based test from SNP-based tests Emily, 2016. Rather than considering multiple SNPs in both gene as part of a joint model, these methods aim at aggregating p-values obtained at the SNP level into a single p-value at a gene level.

## Interaction testing at the SNP level

Let consider a pair of SNPs, $\left(X_{1, j}, X_{2, k}\right)$ where $X_{1, j}$ is the $j^{\text {th }}$ SNP of gene $X_{1}$ and $X_{2, k}$ the $k^{\text {th }}$ SNP of gene $X_{2}$ $\left(1 \leq j \leq m_{1}\right.$ and $\left.1 \leq k \leq m_{2}\right)$. To test for interaction at the SNP level, we used the following Wald statistic:

$$
W_{j k}=\frac{\widehat{\beta_{3}^{j, k}}}{\sigma \widehat{\left(\widehat{\beta_{3}^{j, k}}\right)}}
$$

where $\widehat{\beta_{3}^{j, k}}$ is an estimate of the interaction coefficient $\beta_{3}^{j, k}$ of the following logistic model:

$$
\log \left(\frac{\mathbb{P}\left[Y=1 \mid X_{1, j}=x_{1}, X_{2, k}=x_{2}\right]}{1-\mathbb{P}\left[Y=1 \mid X_{1, j}=x_{1}, X_{2, k}=x_{2}\right]}\right)=\beta_{0}^{j, k}+\beta_{1}^{j, k} x_{1}+\beta_{2}^{j, k} x_{2}+\beta_{3}^{j, k} x_{1} x_{2}
$$

$\widehat{\beta_{3}^{j, k}}$ is obtained by maximizing the likelihood function on the observed data $\mathbf{Y}, \mathbf{X}_{\mathbf{1}}$ and $\mathbf{X}_{\mathbf{2}}$ while $\sigma \widehat{\left(\widehat{\beta_{3}^{j, k}}\right)}$ is calculating by inverting the Hessian of the likelihood. Since the solution of the maximization of the likelihood function does not have a closed form, we compute $W_{j k}$ according to the iteratively reweighted least squares algorithm proposed in the glm function of the stats package R Core Team, 2016 .

To combine the statistics $W_{j k}$ into a single test, Ma et al., 2013 proposed four methods that all account for covariance matrix $\Sigma=\left[\sigma_{(j, k),\left(j^{\prime}, k^{\prime}\right)}\right]_{\substack{ \\j^{\prime}=1 \ldots m_{1} ; k=1 \ldots m_{2}}}$, a $\left(m_{1} \times m_{2}\right) \times\left(m_{1} \times m_{2}\right)$ symmetric matrix where $j^{\prime}=1 \ldots m_{1} ; k^{\prime}=1 \ldots m_{2}$
$\sigma_{(j, k),\left(j^{\prime}, k^{\prime}\right)}=\operatorname{Cov}\left(W_{j k}, W_{j^{\prime}, k^{\prime}}\right)$. As proposed by Emily, 2016, the covariance between $W_{j k}$ and $W_{j^{\prime}, k^{\prime}}$ is estimated by:

$$
\sigma_{(j, k),\left(j^{\prime}, k^{\prime}\right)}=r_{j, j^{\prime}} r_{k, k^{\prime}}
$$

where $r_{j, j^{\prime}}=\frac{p_{j j^{\prime}}-p_{j} p_{j^{\prime}}}{\sqrt{p_{j}\left(1-p_{j}\right) p_{j^{\prime}}\left(1-p_{j^{\prime}}\right)}}$ is the widely used correlation measure between SNP $j$ and SNP $j^{\prime}$, given that $p_{j}$ and $p_{j^{\prime}}$ are the respective allelic frequencies and $p_{j j^{\prime}}$ is the joint allelic frequency Hill and Robertson, 1968.

In the remainder of this section, the four methods: minP (function minP.test, GATES (function gates.test), tTS (function tTS.test) and tProd (function tProd.test) are detailed.

## $3.1 \operatorname{minP}$

The minP test is based on the minimum p-value that is often used to combine p-values of association (see Conneely and Boehnke, 2007). Let $W_{\max }=\max \left|W_{11}\right|, \ldots,\left|W_{m_{1}, m_{2}}\right|$ be the maximum of the absolute observed statistics. The minP is then defined by:

$$
\begin{equation*}
\operatorname{minP}=1-\mathbb{P}\left[\max \left(\left|Z_{1}\right|,\left|Z_{2}\right|, \ldots,\left|Z_{m_{1} m_{2}}\right|\right)<W_{\max }\right] \tag{1}
\end{equation*}
$$

where $\mathbb{Z}=\left(Z_{1}, Z_{2}, \ldots, Z_{m_{1} m_{2}}\right)$ is a random vector that follows a multivariate normal distribution $\mathbb{Z} \sim \mathcal{N}(\mathbf{0}, \Sigma)$.
The computation of Equation (1) requires the calculation of the probability distribution of a multivariate normal random variable. For that purpose, we used the pmvnorm function from the R package mvtnorm Genz and Bretz, 2009.

```
> set.seed(1234)
> minP.test(Y=gene.pair$Y, G1=gene.pair$G1,G2=gene.pair$G2)
    Gene-based interaction based on minP method
data: gene.pair$Y and (gene.pair$G1 , gene.pair$G2)
Wmax = 0.0099241, p-value = 0.1796
alternative hypothesis: true Wmax is greater than 0
sample estimates:
    Wmax
0.009924148
```


### 3.2 GATES

The GATES procedure, proposed by Li et al., 2011, is an extension of the Simes procedure used to assess the gene level association significance. Let $p_{(1)}, \ldots, p_{\left(m_{1} m_{2}\right)}$ be the ascending SNP-SNP interaction $m_{1} \times m_{2}$ p-values, GATES p-value is then defined by

$$
\mathrm{p}_{G A T E S}=\min \left(\frac{m e p_{(1)}}{m e_{(1)}}, \frac{m e p_{(2)}}{m e_{(2)}}, \ldots, \frac{m e p_{\left(m_{1} m_{2}\right)}}{m e_{\left(m_{1} m_{2}\right)}}\right)
$$

where $m_{e}$ is the number of effective tests among the $m_{1} \times m_{2}$ tests and $\left.m e_{( } i\right)$ the number of effective tests among the $i$ most significative tests associated with the lowest order p-values $p_{(1)}, \ldots, p_{(i)}$. The number of effective tests ought to characterize the number of independent tests equivalent to the correlated tests that are really performed and is often used to account for dependence in a multiple testing correction.

Although no formal definition of the number of effective tests has been formulated in the literature, several procedures have been proposed to estimate such number. All methods are based on a transformation of the set of eigenvalues of the SNP covariance matrix assuming that (1) if the SNPs are independent, the number of effective tests is the number of performed, (2) if the absolute value of the correlation between any pair of SNPs is equal to 1 , the number of effective tests is 1 . In the GeneGeneInteR package, four main methods have been implemented and can be chosen by the user with the argument merest: Cheverud-Nyholt method - me.est="ChevNy" Cheverud, 2001, Nyholt, 2004, Keff method -me.est="Keff" Moskvina and Schmidt, 2008, Li and Ji method - me.est="LiJi" Li and Ji, 2005 and Galwey - me.est="Galwey" Galwey, 2009.
> set.seed (1234)
> gates.test(Y=gene.pair\$Y, G1=gene.pair\$G1,G2=gene.pair\$G2,me.est="ChevNy")

Gene-based interaction based on GATES method
data: gene.pair\$Y and (gene.pair\$G1, gene.pair\$G2)
GATES $=0.0099241, \mathrm{p}$-value $=0.2939$
alternative hypothesis: less
sample estimates:
GATES
0.009924148

```
> set.seed(1234)
> gates.test(Y=gene.pair$Y, G1=gene.pair$G1,G2=gene.pair$G2,alpha=0.05,me.est="Keff")
```

            Gene-based interaction based on GATES method
    data: gene.pair\$Y and (gene.pair\$G1, gene.pair\$G2)
GATES $=0.013945, \mathrm{p}$-value $=0.1899$
alternative hypothesis: less
sample estimates:
GATES
0.01394543
> set.seed (1234)
> gates.test(Y=gene.pair\$Y, G1=gene.pair\$G1,G2=gene.pair\$G2,me.est="LiJi")
Gene-based interaction based on GATES method
data: gene.pair\$Y and (gene.pair\$G1, gene.pair\$G2)
GATES $=0.013945, \mathrm{p}$-value $=0.1255$
alternative hypothesis: less
sample estimates:
GATES
0.01394543
> set.seed (1234)
> gates.test(Y=gene.pair\$Y, G1=gene.pair\$G1,G2=gene.pair\$G2,me.est="Galwey")

```
    Gene-based interaction based on GATES method
data: gene.pair$Y and (gene.pair$G1 , gene.pair$G2)
GATES = 0.013945, p-value = 0.1596
alternative hypothesis: less
sample estimates:
    GATES
0.01394543
```


## 3.3 tTS and tProd

tTS and tProd procedures are two truncated tail strength methods that aim at combining signals from all single-SNP p-values less than a predefined cutoff value Jiang et al., 2011. Denoting by $\tau$ the cutoff value, the two truncated p-values are defined as follows Zaykin et al., 2002:

$$
\begin{aligned}
t T S & =\frac{1}{m_{1} m_{2}} \sum_{i=1}^{m_{1} m_{2}} \mathbb{I}\left(p_{(i)}<\tau\right)\left(1-p_{(i)} \frac{m_{1} m_{2}+1}{i}\right) \\
t \text { Prod } & =\prod_{i=1}^{m_{1} m_{2}} p_{i}^{\mathbb{I}\left(p_{i}<\tau\right)}
\end{aligned}
$$

where $\mathbb{I}$ is the indicator function.
When p-values are correlated, the null distribution of $t T S$ and $t$ Prod are unknown. Following the approach proposed by Zaykin et al., 2002, a p-value is obtained in the GeneGeneInteR package by computing an empirical null distribution using Monte-Carlo (MC) simulations. For each MC iteration, an empirical value for $t T S$ (or $t$ Prod) is obtained by simulating a vector of $W_{j k}$ with respect to a multivariate normal distribution with a vector of 0 means and $\widehat{\Sigma}$ as covariance matrix. The empirical p-value is calculated as the proportion of simulated statistics larger than the observed statistic on the "true" set of $W_{j k}$.
tTS and tProd methods have been implemented in the functions tTS.test and tProd.test of the GeneGeneInteR package. Additional to the mandatory Y, $G_{1}$ and $G_{2}$ arguments, these two functions have two optional arguments: tau and n.sim that control the cutoff value and the number of simulations used to estimate the empirical value respectively. The following coding lines give an example of the tTS.test and tProd.test:

```
> set.seed(1234)
> tTS.test(Y=gene.pair$Y, G1=gene.pair$G1,G2=gene.pair$G2,tau=0.5,n.sim=10000)
```

Gene-based interaction based on the Truncated Tail Strength method
data: gene.pair\$Y and (gene.pair\$G1 , gene.pair\$G2)
tTS $=-0.0099127$, tau $=0.5, \mathrm{p}$-value $=0.5104$
alternative hypothesis: less
sample estimates:
tTS
-0.009912706
> set.seed (1234)
> tProd.test(Y=gene.pair\$Y, G1=gene.pair\$G1,G2=gene.pair\$G2,tau=0.05,n.sim=1000)

Gene-based interaction based on the Truncated Product method

```
data: gene.pair$Y and (gene.pair$G1 , gene.pair$G2)
tProd = 0.0001384, tau = 0.05, p-value = 0.265
alternative hypothesis: less
sample estimates:
    tProd
0.0001383965
```


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