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# Software Manual for iOmicsPASS: integrative -Omics Tool for Predictive Analysis with Subnetwork Signatures

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*Version 1.0*

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February 19, 2019

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# 1 Installation

The software requires a **gcc** compiler and makes use of the **boost** library (available at <http://www.boost.org/users/download/>), which is distributed along with the software package under the Boost Software License ([https://www.boost.org/LICENSE\\_1\\_0.txt](https://www.boost.org/LICENSE_1_0.txt)). Additionally, a programming software **R** [1] is required for graphical visualization of the results.

## 1.1 Installing and setting up R

To install **R**, go to the website (<https://cran.r-project.org>) and download the right version (Linux, Mac OS X or Windows) for installation. The latest version of **R** is version 3.4.3. After that, add the directory where **R** is installed to the system **Path** environment variable by specifying the following:

### For Windows users

Navigating through the following:

“My Computer > Control Panel > Systems > Advance system settings > Environment variable” and click on **edit** to add "C:/Program Files/R/R-3.4.3/bin" manually. Or use command line to create/set a variable permanently (as Administrator), use:

```
> setx PATH "C:/Program Files/R/R-3.4.3/bin/"
```

### For Mac OS X/Linux users

Make sure that "usr/local/bin" is already added in **Path**, else type the following in the “Terminal”:

```
> PATH=$PATH:/usr/local/bin/
```

## 1.2 Installing iOmicsPASS

After downloading the zip folder for the software package from github (<https://github.com/cssblab/iOmicsPASS>), uncompress the folder on your local directory. Then, type “make” in the directory to install all the components of the software. Users can add the directory of iOmicsPASS to their PATH variable by specifying:

For example, if the program is installed in "/Users/hiromi/Desktop/",

For BASH/ksh/sh shell users,

```
>PATH=$PATH:/Users/hiromi/Desktop/iOmicsPASS_v1.0/bin/
```

To make these changes permanent,

For BASH shell users,

```
>export PATH=$PATH:/Users/hiromi/Desktop/iOmicsPASS_v1.0/bin/ >> ~/.bash_profile
```

For ksh/sh shell users,

```
>export PATH=$PATH:/Users/hiromi/Desktop/iOmicsPASS_v1.0/bin/ >> ~/.profile
```

### For Mac OS X/Linux users

Full installation of Xcode is required in Mac OS X users and all other instructions are the same as linux.

### For Windows users

Executables for 64-bit Windows are included in the zip folder. Installation of Cygwin [2] is required (download available at <https://www.cygwin.com/>) to run iOmicsPASS and after installation, ensure that the directory is added to the system **Path** environment variable by navigating through the following:

“My Computer > Control Panel > Systems > Advance system settings > Environment variable” and click on **edit** to add "C:/cygwin64/bin" manually. Or use command line to create/set a variable permanently (as Administrator), use:

```
> setx PATH "C:/cygwin64/bin"
```

## 2 Introduction to iOmicsPASS

iOmicsPASS is a powerful bioinformatics tool, developed in C++ language, for network-based data integration and predictive feature selection. The tool integrates multiple -omics datasets utilizing biological network information and identifies important biological interactions in the network as signatures to distinguish between different sample groups (i.e. phenotypes). It is a supervised method where the classification of the samples is known in the training data. The tool can be applied in clinical settings, to better understand the biological mechanisms underlying the differences across groups of samples (e.g. when comparing disease versus non-disease group) in studies with multiple related -omics data.

## 3 Workflow

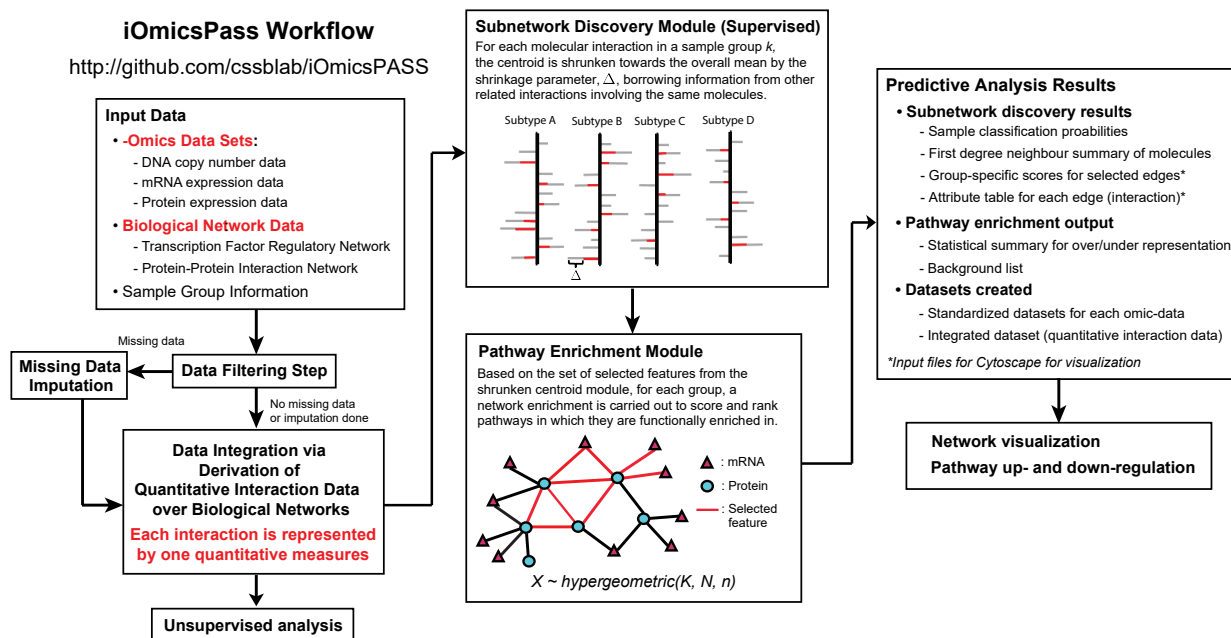


Figure 1: Workflow of iOmicsPASS

Figure 1 above shows the overview of the workflow in iOmicsPASS. The tool first applies data filtering with user-specified criteria on each -omics dataset to ensure that molecules with too many missing data points are removed from the data. Next, if there are still missing data, the user can choose to use the  $K$ -nearest neighbour (KNN) method to impute the data [3]. The multi-omics datasets are then integrated into a single dataset over the user-provided biological network to form an edge-level feature dataset (nodes and edges are used following the nomenclature of network biology). Next, a subnetwork discovery module, adapted

from the Prediction Analysis of Microarray method by Tibshirani’s *et al.* [4], is carried out on the transformed data, shrinking each group’s centroid towards the overall average centroid by using a soft-thresholding method.  $K$ -fold cross-validation is used to optimize the shrinkage parameter, which minimizes the overall misclassification error rate. Finally, a novel method for pathway-level scoring, in a network setting, is carried out for the set of selected features from the previous steps.

Further details regarding the method can be found in Koh *et al.* [5]

### 3.1 Data Filtering Step

To filter out molecules with too many missing data points, each molecule should have non-missing measurements in (1) at least  $x$  number of samples or (2) at least  $x/n$  samples with non-missing data in each sample group, where  $x$  and  $n$  are integers and  $x \leq n$ .

### 3.2 Data Imputation

To impute missing data, weighted  $K$ -nearest neighbors (KNNimpute) method proposed by Schwender *et al.* [6] is used. For every molecule  $i$  with a missing value in sample  $j$ , the algorithm searches for  $K$  other molecules most similar to the given molecule but with a value present in sample  $j$ . A weighted average of the values for the  $K$  closest molecules in the same sample  $j$  is used as an estimate for the missing value, with the weights proportional to the similarity of each of the  $K$  molecules and molecule  $i$ . The similarity is determined by the Euclidean distance, or  $L_2$ -norm.

Specifically, assuming  $\mathcal{L}_K$  is a set consisting of  $K$  molecules with the smallest Euclidean distances to molecule  $i$  with a value present in sample  $j$ . The missing value  $x_{ij}$  is replaced by

$$x_{ij} = \sum_{\ell \in \mathcal{L}_K} w_{i\ell} x_{\ell j} / \sum_{\ell \in \mathcal{L}_K} w_{i\ell}$$

where the weight  $w_{i\ell}$  is the reciprocal of the Euclidean distance between feature  $i$  and feature  $\ell$ .

### 3.3 Integration of Multiple -Omics Datasets

To integrate multiple -omics datasets, we create a measurement of “co-variation” for every pair of interacting molecules in the given biological network. When modelling the expression profiles across samples at the DNA, mRNA and protein level, two types of networks are relevant for linking the information from different molecular levels: protein-protein interaction (PPI) network and transcription factor (TF) regulatory network. The former is a network of physical binding between protein molecules, and the latter is a network between protein molecules of TF genes and mRNA molecules of target genes. In the latter network, DNA copy number can also be incorporated as a normalizing value for mRNA abundance, since the ratio mRNA / DNA copy number can be considered as the “output” of gene transcription. Here we form two types of edges, where we consider the case with DNA copy number and the case without separately.

Each edge,  $e_{i,j,t}$  for  $i=1,\dots, p$ ,  $j=1,\dots, n$  and  $t=1$  or  $2$ , is defined as follows:

(1) when DNA copy number data is not available,

$$\begin{aligned} e_{i,j,1} &= z_{prot_A,j} + z_{mrna_B,j} \\ e_{i,j,2} &= z_{prot_A,j} + z_{prot_B,j} \end{aligned}$$

(2) when DNA copy number data is available,

$$\begin{aligned} e_{i,j,1} &= z_{prot_A,j} + (z_{mrna_B,j} - z_{dna_B,j}) \\ e_{i,j,2} &= z_{prot_A,j} + z_{prot_B,j} \end{aligned}$$

where  $z$  represents the standardized log-scale (base 2) measurement of each molecule in the different -omics datasets,  $p$  denotes the total number of edges,  $n$  denotes the total number of samples, and  $t$  represent the type of data, where  $t=1$  denotes the TF edge and  $t=2$  denotes the PPI edge. For  $t=1$ , gene A is the TF and gene B is the target of A.

Here,  $mrna_B$  and  $dna_B$  are the mRNA expression and DNA copy number of gene B, respectively. It is assumed to be a target of the transcription factor protein  $prot_A$ . Also,  $prot_A$  and  $prot_B$  are assumed to be any two different interacting proteins.

These new values  $\{e\}$  are considered as **co-variation** between two or three related molecules because consistently high or low values of the molecules involved in a biological interaction (PPI or TF regulation) suggests increased or decreased chance of molecular interaction in a given biological sample.

### **Generalizing to other -omics datasets**

The same data transformation approach can also be used to integrate other types of data such as mRNA expression and DNA methylation data (e.g. DNA methylation array with probe sets located in various genomic regions). Similar to protein-mRNA integration, a gene-to-gene interaction network and a network of methylation probes to their nearest gene(s) can be used for integration. It is commonly reported in literature [7, 8, 9] that methylation probes are usually negatively correlated with the gene target's expression values if the methylation site is located in the upstream of transcription start site (TSS) of a gene, and are positively correlated if the site is located within a gene body. In this case, we multiply a signed constant (-1 or 1) to the standardized measurements and allow for positive and negative interactions, as directed by the user.

The edge data are calculated as follows:

$$e_{i,j,1} = z_{gene_A,j} + sign(z_{methylprobe_B,j}) * z_{methylprobe_B,j}.$$

The same logic can be applied for the integration of negative interactions for microRNAs and protein and mRNA molecules of their target genes based on a target scan map of conserved sequences of miRNAs and target genes.

## **3.4 Subnetwork Discovery Module**

In this module, the integrated data is put into a shrunken-centroid algorithm that we modified from the traditional PAM methodology to identify edges in a network that is predictive of the sample groups. This

involves testing and training of data using cross-validations to build the best model that can be used to predict group membership of samples. And by doing so, we also obtain the set of edges or biological interactions in the overall network constructed, that best characterizes each of the sample groups that are called **signatures**.

### 3.4.1 Modifications to PAM method for predictive analysis

Tibshirani’s *et al.* [4] developed a method for identifying important gene signatures across multiple subtypes of cancers using gene expression microarray data, called Prediction of Microarray (PAM) which is a nearest shrunken centroid method (NSC). The method computes a group centroid for each feature (gene) with a soft-thresholding method, and the shrinkage parameter is optimized using cross-validation. However, the PAM method was designed for a single -omic dataset and the method was shown to be biased towards subgroups with larger sample sizes [10, 11]. As part of i0micsPASS, we made changes to the PAM method to account for the network-based data integration. The modifications are as follows:

#### (1) Computation of centroids for each feature (edge-level data)

The overall centroid for feature  $i$ , or interaction  $i$ , is weighted by sample group size and is calculated as:

$$\bar{x}_i = \frac{1}{K} \sum_{k=1}^K \bar{x}_{ik} = \frac{1}{K} \sum_{k=1}^K \left( \frac{1}{n_k} \sum_{j \in C_k} x_{ij} \right)$$

where  $C_k$  represents the set of indices of the  $n_k$  samples in group  $k$  and  $K$  represents the number of sample groups.

#### (2) Computation of test statistics for features

In PAM,  $d_{ik}$  is defined to be the  $t$  statistics for each gene  $i$ , comparing sample group  $k$  to the overall centroid:

$$d_{ik} = \frac{\bar{x}_{ik} - \bar{x}_i}{m_k \cdot (s_i + s_0)}$$

where  $m_k = \sqrt{1/n_k + 1/n}$  and  $s_i$  is the pooled within-class standard deviation of feature  $i$  and  $s_0$  is a positive constant, the median value of  $s_i$  across all features.

Here, we add an additional term in the score to reward or penalize  $d_{ik}$  depending on the quantitative levels of the “neighbor” edges that share common molecules. We define  $d_{ik}^*$  as the new test statistics for each edge  $i$ , comparing sample group  $k$  to the overall centroid:

$$d_{ik}^* = d_{ik} + \left( \psi_k * \frac{|N_{e_{i,1}}| \sum_{s \in N_{e_{i,1}}} d_{sk} + |N_{e_{i,2}}| \sum_{r \in N_{e_{i,2}}} d_{rk}}{|N_{e_i}|} \right)$$

where  $N_{e_i}$  represents the set of edges which are connected to at least one of the nodes of edge  $e_i$ .  $N_{e_i}$  can be further partitioned into two subsets,  $N_{e_{i,1}}$  and  $N_{e_{i,2}}$ , representing the set of TF and PPI edges, respectively. Furthermore, we set

$$\psi_k = \frac{2e^{5(p_{ik}-0.5)}}{1 + e^{5(p_{ik}-0.5)}}$$

where

$$p_{ik} = \frac{|\sum_{j \in N_{e_i}, i \neq j} \text{sign}(d_{ik}) = \text{sign}(d_{jk})|}{|N_{e_i}|}.$$

Here,  $\psi_k$  acts as a multiplicative factor based on the proportion of agreement,  $p_k$ , between the edge and its direct neighbor edges sharing at least one node of the edge  $i$  in sample group  $k$ .

**Note.** Since the set of neighbor edges sharing node(s) of a given edge is invariant across samples,  $e_{ijt}$  is referred to as  $e_i$  ( $e_{i1}$  for TF edges and  $e_{i2}$  for PPI edges) below.

### (3) Group-specific thresholds for soft-thresholding

For each sample group  $k$ , a group-specific threshold  $\Delta_k$  is derived from the shrinkage parameter,  $\Delta$ .

$$\Delta_k = \frac{\Delta}{\Delta_{max}} * \max d_{ik}$$

where

$$\Delta_{max} = \frac{1}{K} \sum_{k=1}^K \max d_{ik}$$

$\Delta$  is defined to be over a grid of 30 equally-spaced values arranged in an increasing order from zero to some constant (i.e.  $\Delta \in \{0, \Delta_1, \Delta_2, \dots, \Delta_{max}\}$ ). Using a soft-thresholding method, each feature's distance measure  $d_{ik}$  is re-computed by reducing it incrementally by an absolute shrinkage amount and it will be set to zero if it falls below zero:

$$d'_{ik} = \text{sign}(d_{ik}^*)(|d_{ik}^*| - \Delta_k)_+$$

The optimal value for  $\Delta$  is chosen based on  $K$ -fold cross-validation, to yield a set of features, or subnetwork signatures that can best classify samples to their respective groups.

#### 3.4.2 Cross-validations

By default, a 10-fold cross-validation is used to estimate the shrinkage parameter for soft-thresholding. The data is first split into 10 mutually exclusive parts where each part is used as test dataset and the rest of the 9 parts are used as training datasets. The training dataset is used to build a discriminant model that separates the sample groups based on a certain  $\Delta$  value. Then the model is used to predict the membership of the test samples and the mis-classification error is recorded. This is repeated over a grid of increasing values for  $\Delta$  starting from 0 to a value large enough that all the features in the model are reduced towards the group centroids (i.e.  $d'_{ik}$  values become zero) and this is known as soft-thresholding method. The process is repeated for each of the 10 folds and the mis-classification error recorded will be averaged to give the overall mis-classification error.

The test samples will be classified to the nearest shrunken centroid of each group. Suppose a test sample with expression values  $x^* = (x_1^*, x_2^*, \dots, x_p^*)$ , the discriminant score for class  $k$  is defined to be:

$$\delta_k(x^*) = \sum_{i=1}^p \frac{(x_i^* - \bar{x}'_{ik})^2}{(s_i + s_0)^2} - 2\log(\pi_k)$$



where  $\pi_k$  is the  $k^{th}$  class prior probability and estimated using an equal prior  $\pi_k = 1/K$ .

We then assign each of the test sample to the class with the smallest discriminant score, using the classification rule:  $C(x^*) = \ell$  where  $\delta_\ell(x^*) = \min_k \delta_k(x^*)$ . Using the discriminant scores, we can then construct the estimates of the class probabilities as follows:

$$\hat{p}_k(x^*) = \frac{e^{-\frac{1}{2}\delta_k(x^*)}}{\sum_{\ell=1}^K e^{-\frac{1}{2}\delta_\ell(x^*)}}.$$

### 3.5 Pathway Enrichment Module

From the previous step, a set of edges with non-zero centroids are obtained. These edges can be used to perform enrichment of biological pathways in each of the subgroup. The edges are first separated into two groups with positive and negative  $d'_{ik}$  scores for each subgroup. Edges which have been shrunk towards zero (i.e.  $d'_{ik} = 0$ ) are removed. Then, hypergeometric test will be applied to compute the probability of over-representation in a particular pathway, separately for each subgroup.

Given a graph  $G = (V, E)$  where  $V$  is the set of all possible nodes (i.e. mRNA and Proteins available in the data) and  $E$  is the set of all possible edges (i.e. TF and PPI interactions from network information) in the graph. We can construct smaller subgraphs,  $G_p = (V_p, E_p)$  to represent every pathway where  $V_p$  (i.e.  $V_p \subset V$ ) is the set of nodes representing all genes in pathway  $p$  and  $E_p$  (i.e.  $E_p \subset E$ ) is the set of edges that exist in the graph. Using the set of edges in the enrichment list for each sample group  $k$ , we create an induced subgraph of graph  $G_p$ ,  $G_k = (V_k, E_k)$ , where  $V_k$  (i.e.  $V_k \subseteq V_p$ ) is the set of nodes in the enrichment list that represent all genes in pathway  $p$ , and  $E_k$  (i.e.  $E_k \subseteq E_p$ ) is the set of edges in the enrichment list that exist in the subgraph.

To find the probability of over-representation of edges in a particular pathway, we perform a hypergeometric test. Let  $X$  be a random variable representing the number of edges in a graph and define the following:

$$X \sim \text{Hypergeometric}(K, N, n)$$

$$Pr(\text{overrepresentation}) = Pr(X > x) = 1 - Pr(X \leq x)$$

where  $|E_k| = K$  is the number of edges in the enrichment list,  $|E| = N$  is the total number of possible TF and PPI edges, and  $n$  is the number edges that can be constructed with the vertices representing genes that belong to the pathway of interest.

## 4 Structuring of Input Datasets

iOmicsPASS takes several types of input files and the following sections give an overview of the required structure of the input datasets. All input files should be formatted in tab-delimited format. The labels in the header do not matter but the ordering of the information in each column for the different input files should be strictly adhered to the format as described below.

### 4.1 Molecular Data

The molecular data should be in the format of a  $p$  by  $n$  matrix, where the rows represent the  $p$  molecules and the columns represent the samples. The first column in the data is the gene/protein identifier, followed by quantitative values in each sample. The column headers for the samples should be unique and have to correspond to the sample IDs provided by user in the sample group information.

Table 1: Example of a molecular dataset structure

Gene_identifier	Sample_1	Sample_2	...	Sample_n
<i>gene</i> <sub>1</sub>	1.032	1.423	...	0.894
<i>gene</i> <sub>2</sub>	2.355	0.983	...	1.188
<i>gene</i> <sub>3</sub>	0.783	1.216	...	0.730
<i>gene</i> <sub>4</sub>	2.892	1.822	...	0.741
⋮	⋮	⋮	⋮	⋮
<i>gene</i> <sub><math>p</math></sub>	0.283	1.024	...	0.762

**Note.** The ordering of the samples does not matter and data will only be extracted from samples that have a sample ID provided in the sample group information. The type of gene identifiers should be uniform across the different -omics datasets. For example, if gene symbols are used in RNA-level data, it should also be used in Protein-level data.

### 4.2 Sample Group Information

The sample group information should contain two columns. The first column is the sample identifiers/IDs and the second column is the sample group information (class).

Table 2: Example of a Sample Information structure

Sample_ID	Classification
Sample_1	Class_1
Sample_2	Class_3
Sample_3	Class_2
Sample_4	Class_1
⋮	⋮
Sample_n	Class_k

### 4.3 Network Data

Up to two types of network files can be supplied by the user in the current version. One of the network files will be used to link features from the Protein-level input data to the RNA-level input data. The other network file will be used to link features within the Protein-level input data.

In the TF network file, there should be two columns containing headers where each row shows the interaction between two molecules: the molecule in the first column has a directed edge pointing or targeting the corresponding molecule in the second column (see Table 3A).

In the PPI network file, there should be two columns containing headers where each row shows the interaction between two features: the node in the first column has an undirected edge with the corresponding node in the second column (see Table 3B).

For implementing both positive and negative interactions in a single network, a third column should be included in the network file with values 1 or -1, to denote positive and negative interactions, respectively (see Table 3C).

Table 3A: Example of a TF network file structure

<b>TF</b>	<b>TF_target</b>
<i>Protein<sub>a</sub></i>	<i>mRNA<sub>1</sub></i>
<i>Protein<sub>a</sub></i>	<i>mRNA<sub>2</sub></i>
<i>Protein<sub>b</sub></i>	<i>mRNA<sub>3</sub></i>
<i>Protein<sub>b</sub></i>	<i>mRNA<sub>4</sub></i>
<i>Protein<sub>c</sub></i>	<i>mRNA<sub>5</sub></i>
$\vdots$	$\vdots$

Table 3B: Example of a PPI network file structure

<b>Protein_A</b>	<b>Protein_B</b>
<i>Protein<sub>a</sub></i>	<i>Protein<sub>b</sub></i>
<i>Protein<sub>a</sub></i>	<i>Protein<sub>c</sub></i>
<i>Protein<sub>b</sub></i>	<i>Protein<sub>d</sub></i>
<i>Protein<sub>c</sub></i>	<i>Protein<sub>d</sub></i>
<i>Protein<sub>c</sub></i>	<i>Protein<sub>e</sub></i>
$\vdots$	$\vdots$

Table 3C: Example of a network file with signed interactions

<b>Gene</b>	<b>MethylationProbe</b>	<b>InteractionSign</b>
<i>gene<sub>a</sub></i>	<i>methyprobe<sub>1</sub></i>	1
<i>gene<sub>a</sub></i>	<i>methyprobe<sub>2</sub></i>	-1
<i>gene<sub>b</sub></i>	<i>methyprobe<sub>3</sub></i>	-1
<i>gene<sub>b</sub></i>	<i>methyprobe<sub>4</sub></i>	1
<i>gene<sub>c</sub></i>	<i>methyprobe<sub>5</sub></i>	1
$\vdots$	$\vdots$	$\vdots$

## 4.4 Pathway Information

The pathway information file should have three columns containing headers. The first is the gene identifier, the second column is the pathway identifier that the gene belongs to and the third column denotes the pathway information (e.g. name of pathway). The type of gene identifier used should be consistent with that used in the input data. For example, if gene symbols are used in the input for molecular datasets, then it should also be used here in the pathway information.

Table 4: Example of a Pathway module file structure

Gene_Symbol	Pathway_ID	Pathway_Function
SOX9	GO:0042981	regulation of apoptotic process
MAP3K8	GO:0042981	regulation of apoptotic process
ALX4	GO:0042981	regulation of apoptotic process
UBC	GO:0042981	regulation of apoptotic process
BCL6	GO:0000902	cell morphogenesis
FRYL	GO:0000902	cell morphogenesis
SOX6	GO:0000902	cell morphogenesis
⋮	⋮	⋮

## 5 Specification for Input Parameter file

The table below (Table 5) gives a short description of each input parameter offered in iOmicsPASS.

Table 5: Description of input parameters for iOmicsPASS

Parameter Label	Variable Type	Description
<i>Input data files</i>		
DNA_FILE	<i>string</i>	Filename of DNA copy number dataset (optional).
RNA_FILE	<i>string</i>	Filename of RNA-level expression dataset.
PROT_FILE	<i>string</i>	Filename of Protein-level expression dataset.
PPI_NETWORK	<i>string</i>	Filename of Protein-Protein interaction network file.
TF_NETWORK	<i>string</i>	Filename of Transcription-factor regulatory network file.
MODULE_FILE	<i>string</i>	Filename of pathway information for enrichment.
SUBTYPE_FILE	<i>string</i>	Filename of group information.
<i>Parameters for data manipulation and filtering step</i>		
LOG_TRANSFORM_DNA	<i>boolean</i> <true/false>	If true, the data will be treated as intensities (before log) and log (base 2) transformation will be carried out. If false, no log-transformation will be carried out.

*Continued on next page*

Table 5 – continued from previous page

Parameter Label	Variable Type	Description
LOG_TRANSFORM_RNA	<i>boolean</i> <true/false>	If true, the data will be treated as intensities (before log) and log (base 2) transformation will be carried out. If false, data will be assumed to be log-transformed (default is true).
LOG_TRANSFORM_PROT	<i>boolean</i> <true/false>	If true, the data will be treated as intensities (before log) and log (base 2) transformation will be carried out. If false, data will be assumed to be log-transformed (default is true).
ZTRANS_DNA	<i>boolean</i> <true/false>	Whether or not to perform standardization to the DNA-level data before integration (default is true).
ZTRANS_RNA	<i>boolean</i> <true/false>	Whether or not to log-transform the RNA-level data before integration (default is true).
ZTRANS_PROT	<i>boolean</i> <true/false>	Whether or not to log-transform the Protein-level data before integration (default is true).
MIN_OBS	<i>integer</i>	Minimum number of samples in each sample group with complete data that is used in the analysis (default is 1).
MIN_PROP	<i>double</i> [0-1]	Minimum proportion of samples in each sample group with complete data that is used in the analysis (default is 0.5).
KNN_IMPUTE	<i>boolean</i> <true/false>	Whether or not to perform K-nearest neighbor imputation. Should be set to true if dataset has missing cells (default is true).
MAX_BLOCKSIZE	<i>integer</i>	Maximum number of samples used to infer the imputation in KNN (default is 1500).
<i>Parameters for iOmicsPASS</i>		
ANALYSE_DNA	<i>boolean</i> <true/false>	Whether or not to include DNA in the integration analysis. If true, a filename has to be provided in DNA_FILE.
INTERACT_SIGN	<i>boolean</i> <true/false>	Whether or not to account for different types of interactions in network. If true, a third column with values 1 or -1 should be included in TF_NETWORK (default is false).
CROSS_VALIDATION	<i>boolean</i> <true/false>	Whether or not to carry out cross-validation to select optimal threshold cut-off. If true, the minimum threshold will be automatically selected and used as a cut-off. If false, a threshold value should be provided in MIN_THRES.
CV_FOLD	<i>integer</i>	Number for $K$ in $K$ -fold cross-validation (default is 10).
MIN_THRES	<i>double</i>	Threshold to be used as the final cut-off in scoring algorithm. If not specified, this value will be automatically selected to be the threshold at which the lowest misclassification error occurs from the cross-validations.
<i>Specifications for subnetwork enrichment module</i>		
BACKGROUND_PROP	<i>double</i>	Minimum proportion of genes in the pathway that are also in the background list (default = 0.5).
MINBG_SIZE	<i>integer</i>	Minimum number of edges constructed from the set of genes in a given pathway (default is 1).
MINSIG_SIZE	<i>integer</i>	Minimum number of selected edges from the network constructed in a given pathway to be reported (default is 3).

#### Choice of threshold through cross-validation:

The tool, by default, will use the threshold that yields the smallest misclassification error to select features if no threshold is specified. At times, this may not be the most desirable method. For instance, there are

many thresholds with similar misclassification error rates, yet the numerically optimal threshold leads to selection of too many features (i.e. there is an alternative threshold with far sparser and more interpretable size of networks).

A separate R-code (`PerformancePlot.R`) is provided in the tool to assist visualization of the misclassification error plot for the entire range of possible thresholds. In the plot, a line indicating one standard deviation above the minimum misclassification error is drawn. Users can select a threshold that produces a more sparse network which maintains the core constituents of predictive network.

More generally, the user can then make an informed decision as to where to choose the optimal threshold where the trade-off between misclassification error rate and the number of selected features is balanced. Then, users can rerun `iOmicsPASS` by turning the option `CROSS_VALIDATION` off and specifying the preferred threshold in `MIN_THRES`.

## 6 Illustration of iOmicsPASS

To illustrate the use of `iOmicsPASS`, we utilize the breast cancer (BRCA) dataset from the Cancer Genome Atlas (TCGA) [12]. The gene expression data was quantified using RNA sequencing in 1,098 tumor samples collected from patients and quantification of proteins using iTRAQ was carried out on a subset of 105 breast cancer patients by the Clinical Proteomic Tumor Analysis Consortium (CPTAC) [13]. Of those, 103 subjects had subtype classification using mRNA-based PAM50 signature [14]. into 4 breast cancer subtypes: Basal-like, Her2-enriched, Luminal A and Luminal B subtype. By performing this analysis, we aim to identify subnetwork signatures of transcriptional regulation network and protein-protein interaction network which are predictive of each breast cancer subtype.

To reduce computational time, we randomly selected 1000 transcription factor proteins and 3000 mRNAs from the gene targets of the selected transcription factors. Then, a smaller subset of the TF and PPI network was derived to make up the datasets in the example folders. Out of the 103 individuals with group information, 24 were classified as Basal-like, 18 were Her2-enriched, 24 were Luminal A and 32 were Luminal B subtype.

Then, `iOmicsPASS` was executed to integrate the proteomics, mRNA and CNV datasets using the input parameters shown in Figure 2. Here, we performed log-transformation on the mRNA data as the values are fragments per kilobase of transcript per million mapped reads (FPKM) and `KNN` imputation has to be set to `true` as there are missing values in two of the datasets. We also require that for each gene/protein in the input data should have at least 10 or 80% (which ever is larger) non-missing observations across the samples in each sample group. For parameters in the enrichment module, we will require pathways with at least 50% of the genes in the background list and report pathways with a size of 3 edges and have at least 1 enriched edge.

In the directory of the example folder, type the following command to run the tool:

```
> ../bin/iOmicsPASS <input_example>
```

If running on Windows, users can use **Command Prompt** and go to the directory "Windows Binary" to use the executable (iOmicsPASS.exe). Note that another input parameter file —input\_windows, slightly modified from input\_example, was created as the directories of the input files changed since iOmicsPASS.exe has to be executed within "Windows Binary" folder.

Then, type the following command to run the tool:

```
> iOmicsPASS.exe <input_windows>
```

**Note:** The lines in the input parameter file commented out by hash # symbol will not be read in. The input parameter and the data files should be placed are in the same working directory. Otherwise, include the full path to the files in the input parameter.

```

#####
### input parameter file ###
#####

## input files ##
DNA_FILE = CNV_BRCA_example.txt
RNA_FILE = mRNA_BRCA_example.txt
PROTEIN_FILE = Proteomics_BRCA_example.txt

## Network files ##
PPI_NETWORK = PPInetwork_example.txt
TF_NETWORK = TFnetwork_example.txt
MODULE_FILE = PathwayEnrichment_example.txt

## subtyping file ##
SUBTYPE_FILE = BRCA_groupinfo_example.txt

## standardardisation ##
ZTRANS_DNA = true
ZTRANS_RNA = true
ZTRANS_PROT = true

## whether or not to perform log(base2) transformation ##
LOG_TRANSFORM_DNA = false
LOG_TRANSFORM_RNA = true
LOG_TRANSFORM_PROT = false

## parameters within each group ##
MIN_OBS = 10
MIN_PROP = 0.8
### only one of the two has to be specified, MIN_PROP = 0.5 and MIN_OBS=1 by default ###
### if both specified, apply both simultaneously ###

## Apply KNN to impute for missing values ##
KNN_IMPUTE = true
MAX_BLOCKSIZE = 1500

## Specify if there is different directions in the type of interaction edges ##
INTERACT_SIGN = false

ANALYSE_DNA = true
## if true, filename should be provided for DNA_file
## if protein-lvl data is not available, one can also replace protein with rna-level data

## parameter for shrunken centroid ##
CROSS_VALIDATION = true
CV_FOLD = 10
#MIN_THRES

## parameter for subnetwork enrichment ##
BACKGROUND_PROP = 0.5
MINBG_SIZE = 3
MINSIG_SIZE = 1

```

**Input data files**

**Data manipulation and filtering**

**Specifications for iOmicsPASS**

**Specifications for Enrichment module**

Get Help WriteOut Read File Prev Page Cut Text Cur Pos  
Exit Justify Where Is Next Page UnCut Text To Spell

Figure 2: Screen shot of input parameter

## 7 Output from iOmicsPASS

Executing iOmicsPASS may take some time depending on the size of the input data (number of molecules) and the density of the network data. The runtime is the longest when the network density is very high, i.e. many interactions share common nodes (molecules). During the runtime, the progress of the software will be printed on the screen. Figure 3 shows an example of a log file when executing the tool on an example dataset.

The total runtime of iOmicsPASS on the example dataset was about 6 minutes. A total of 14,888 edges with quantitative data were constructed and 6,228 features were selected across the four subtypes by the cross-validation.

**Notes:** The number of features (edges) selected for each run may vary as the software uses a random number generator to perform the  $K$ -fold cross-validation. However, the key constituents of the predictive subnetworks should remain stable.

Running iOmicsPASS will yield three types of output files:

- 1) **SubnetworkDiscovery results** - scores and attributes of the selected features, classification probabilities and network summary of nodes on the network.
- 2) **Pathway enrichment** - statistical summary of the enrichment in pathways of the selected features and the list of all genes/proteins used in the enrichment.
- 3) **Datasets created** - standardized datasets and the network-level data.

### 7.1 Subnetwork Discovery results

Four files are generated to summarize the information in the overall network formed by the selected edges predictive of each sample group and report the classification probabilities on the samples based on the set of selected edges:

#### 7.1.1 EdgesSelected\_minThres.txt

Edges included in this output are the predictive subnetwork signatures in at least one of the groups. This file can be read into Cytoscape directly as a table of interactions for network visualization.

- **Edge** : Concatenated string of the two molecules involved in the interaction.
- **NodeA** : Molecule represented on the Node A, interacting with Node B.
- **NodeB** : Molecule represented on the Node B, interacting with Node A.
- **InteractionType** : Type of interaction: <tf> for the co-expression of the TF protein molecule in Node A and the target mRNA molecule in Node B in the TF network; or <ppi> for the co-expression of the protein molecule in Node A and the protein molecule in Node B in the PPI network.
- **XXX\_dikNew** : The test statistics of the edge for the sample group as a measure of the predictiveness. A large absolute value of this score indicates that the edge is highly predictive of the sample group.



```
example — iOmicsPASS input_example — 120x50
Hiromis-MacBook:example hiromi$ ~/Desktop/iOmicsPASS_v1.0/bin/iOmicsPASS input_example

Initiating iOmicsPASS...

log DNA: false
log RNA: true
log prot: false

Applying the following filter (specified by user) for integrity of genes across subtypes:
Minimum number of non-missing observations for each gene within each subtype: 10
Minimum proportion of non-missing observations for each gene within each subtype: 0.8

Reading in the subtype information file....
Number of subtypes found is 4 with 103 number of subjects and the subtypes are the following:
Basal-like: 24
Luminal A: 29
HER2-enriched: 18
Luminal B: 32

Reading in the DNA-level file.....
Reading in DNA.....3000
There are 3000 genes in the DNA-level file.

Reading in the RNA-level file.....
Reading in RNA.....3000
There are 2853 genes in the RNA-level file.

Reading in the Protein-level file.....
Reading in Protein.....1000
There are 813 genes in the Protein-level file.

Reading in the TF network file.....
Reading in the PPI network file.....
There are a total of 14888 edges constructed from both TF and PPI network file.

Reading in the Pathway module file....
There are a total of 14598 number of pathways.

All data files have been successfully read into iOmicsPASS!

Proceeding to next step...

There are 103 subjects with subtype information and 3000 genes after filtering in the DNA-level file
There are 103 subjects with subtype information and 2853 genes after filtering in the RNA-level file
There are 103 subjects with subtype information and 813 genes after filtering in the Protein-level file

KNN imputation will be carried out on RNA and Protein datasets.

Imputing RNA-level data...
=====
Total number of values imputed: 482
Took : 1.41671 seconds.

Imputing Protein-level data...
=====
Total number of values imputed: 821
Took : 1.04674 seconds.

Number of common subjects with all 3 types of data is: 103
There are a total of 2669 TF targets where PSME3 has the most TF targetor (parent nodes) of 63.

Generating output for information on the first-degree neighbours of all features in the data...

There is a total of 3652 features with TF/PPI network interactions and 14 features with no interactions and are removed.
ESR1_prot is the feature with the most first-degree neighbours, 76 RNA targets and 133 PROT tfs/interactors.

Carrying out shrunken centroid module on the interaction edges...
Maximum Dij (average) in mRNA and PROT data is : 6.59916
Class-specific thresholds are: 8.28849 4.67346 7.35558 6.07911
Carrying out cross-validation...(This may take a while)
Number of samples used for crossvalidation in each subtype:
Basal-like: 24
HER2-enriched: 18
Luminal A: 29
Luminal B: 32

Carrying out cross-validation on fold.....10
At the minimum threshold, a total of 6228 edges survived across the subtypes and within each group, number of edges survived that are:
Basal-like: 2399
HER2-enriched: 1804
Luminal A: 3767
Luminal B: 1484
Calculating class probabilities on samples...
The overall misclassification error based on the model trained in cross-validation is 32%.

Starting Network enrichment on surviving edges for each subtype...
Pathway Enrichment Module
The program has finished running!
Time to completion of iOmicsPASS is : 6 minutes
```

Reading in the Data

Missing Data Imputation

Number of samples common across datasets integrated

Subnetwork Discovery module

Different thresholds were applied to each subtype in the shrinkage module

Progress update on K<sup>th</sup> fold of CV

Number of features selected using the optimal threshold

Overall misclassification error rate

Pathway Enrichment Module

Total run-time

Figure 3: Screen shot of executing iOmicsPASS on the example datasets, with annotations.

- **XXX\_sigEdge** : Type of significance of the selected edge: <died> if the edge has a test statistics of 0; <ppi> if the test statistics is not zero and the type of interaction is PPI; <tf> if the test statistics is not zero and the type of interaction is TF.
- **XXX\_dir** : The direction of the selected edge: <died> if the edge has a test statistics of 0; <up> if the test statistics is positive; <down> if the test statistics is negative.
- **XXX\_absdiknew** : The absolute value of the test statistics of the edge for the sample group as a measure of the predictiveness.

### 7.1.2 AttributesTable.txt

This is the attribute table for the nodes in the set of selected edges. This file can be used as an attribute table in Cytoscape.

- **Node** : Identifiers used for nodes.
- **Gene** : Gene identifier provided by user.
- **Type** : Molecule type: <mRNA> if the molecule is a mRNA; <prot> if the molecule is a protein.
- **XXX\_surv** : “Survival” or selection status of the edge: <died> if the edge has a test statistics of 0; <mRNA> if the test statistics is not zero and the molecule is a mRNA; <prot> if the test statistics is not zero and the molecule is a protein.

### 7.1.3 Node\_Neighbors.txt

This file gives a summary of number of “direct neighbors” of each node in the integrated network constructed.

- **Node** : Identifiers used for nodes.
- **Gene** : The gene identifier provided by user.
- **Type** : Type of molecule: <mRNA> if the molecule is a mRNA; <prot> if the molecule is a protein.
- **Neighbor\_inData** : A concatenated string of molecules that have an edge with the current molecule (direct neighbors).
- **NumNeigh\_inData** : Number of nodes connected with the current molecule (by an edge).

### 7.1.4 SampleClass\_Probabilities.txt

This file gives the classification probabilities (calculated based on discriminant scores) of each sample to the different sample groups based on the set of selected features obtained from cross-validation.

- **Subject** : Sample ID
- **Prob\_XXX** : Classification probability: Probability of the sample belonging to each of the sample group.
- **TrueClass** : The sample group that the sample belongs to.
- **PredictedClass** : The sample group that the sample would be assigned to, given the highest classification probability. <tied> is reported if there are more than one highest classification probability score.

## 7.2 Pathway Enrichment

For each sample group, pathway enrichment is tested on the set of up-regulated and down-regulated subnetworks, separately, and a statistical summary of the enrichment is reported. Here, the test of enrichment is performed accounting for the fact that the data are network edges. For each pathway, a smaller network is constructed based on the set of gene identifiers in the pathway and hypergeometric test is used to compute the statistical significance of enrichment for the selected edges (up and down separately).

For example, in the TCGA BRCA dataset, there were 4 cancer subtypes (Basal-like, Her2-enriched, Luminal A and Luminal B) specified, hence there is a total of 8 enrichment files generated based on the set of features selected within each of the subtype.

e.g. **Basal-like\_Enrichment\_up.txt**

- **Pathway** : The pathway identifier provided by user in pathway module file.
- **PathAnnotation** : The annotation of the pathway provided by user in pathway module file.
- **HypergeoPval** : Hypergeometric  $P$ -value for functions in the up- and down-regulated edges.
- **NumGene\_inPathway** : The number of genes in the pathway.
- **NumGenes\_inbg** : The number of genes that are in the pathway and present in the input data, used in the analyses (background list).
- **PropPathway** : The proportion of genes in the pathway that is represented in the background list.
- **NumEdgesformed\_inbg** : The number of edges formed from the set of genes/proteins in the pathway.
- **Enriched\_Edgesize** : The number of edges that are in the enrichment list that belong to the set of edges constructed in the pathway

## 7.3 Datasets Reported by iOmicsPASS

For each of the molecular-level, a dataset will be generated after data-filtering, log-transformation (if required) and standardization (i.e. **Ztransform\_XXX.txt**). If imputation was required, the data will first undergo filtering and log-transformation, followed by imputation where the output will also be generated (i.e. **imputedData\_XXX.txt**), before standardization occurs.

## 8 Visualisation of Analysis Output

Along with the package, we provide a supplementary R-script in the folder **Rcodes**, to aid visualization for evaluating the performance of the cross-validation. Users can use this plot to pick a threshold with good trade-off between the mean misclassification error and number of features selected. Then, re-specify the threshold (i.e. by entering the value in **MIN\_THRES**) in the input parameter and run **iOmicsPASS** again, without carrying out cross-validation (i.e. set **CROSS\_VALIDATION** = false). This is useful especially when

the optimal threshold picked out by the tool by default, results in too many features being selected.

After running `iOmicsPASS`, enter the following command in the same directory where the results are generated (i.e. `example/Windows Binary folder`):

```
> R CMD BATCH ../Rcodes/PerformancePlot.R
```

## 8.1 CVplot\_Penalty.pdf

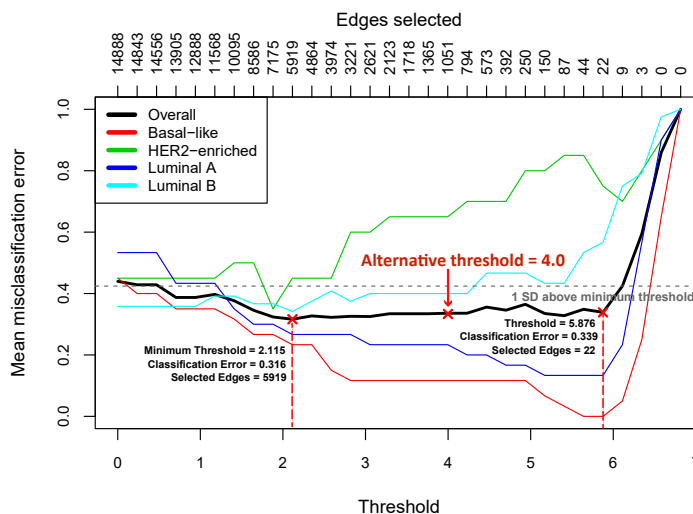


Figure 4: Performance Plot of  $K$ -fold cross-validations

A plot similar to the figure above will be produced. The black colored line shows the overall misclassification error rates and other colored lines show the group-specific misclassification error rates. We observed that, as the threshold increases, the overall misclassification error rate decreases slightly before plateauing and eventually rising to 1 as the number of features selected approaches 0. In this example, the threshold 2.115 is selected as the optimal threshold with the lowest error rate of 31.6%, which results in about 5,919 features selected.

**Note:** The number of features that are selected in the actual data (i.e. 6,228 edges) is slightly different from the number reported in the performance plot (i.e. 5,919 edges) as the latter is calculated using the mean of the features selected from each of the training datasets in  $K$ -fold cross-validations.

A dotted-line is also shown on the plot to illustrate the range of thresholds that keep the mis-classification error rates below one standard deviation (SD) from the minimum error. This enables the user to pick another threshold that provides good trade-off between number of edges selected and the error rate. For example in figure 4, thresholds ranging between 2.1 to 6.0 results in errors within one SD away from the minimum error of 31.6%. At the current threshold that minimizes the error, the number of edges selected may be still too large and to reduce this number, hence making the edges selected more specific, an alternative threshold should be considered. One could pick a threshold of 4.0 instead and the number of edges selected would be reduced to about 1,000 and at the same time, not increasing the error by too much and keeping it controlled at about 32%. The user can then re-run `iOmicsPASS` without carrying out cross-validation (saving computation time) and specifying a threshold in the input parameter file. One can also replace the names

of the input files with the standardized versions (i.e. **Ztransform\_XXX.txt**) produced in the previous run and turn off all options for standardization and log-transformation. (see Figure 5)

```

#####
### input parameter file ###
#####

## input files ##
DNA_FILE = Ztransform_DNA.txt
RNA_FILE = Ztransform_RNA.txt
PROTEIN_FILE = Ztransform_PROT.txt

## Network files ##
PPI_NETWORK = PPInetwork_example.txt
TF_NETWORK = TFnetwork_example.txt
MODULE_FILE = PathwayEnrichment_example.txt

## subtyping file ##
SUBTYPE_FILE = BRCA_groupinfo_example.txt

## standardisation ##
ZTRANS_DNA = false
ZTRANS_RNA = false
ZTRANS_PROT = false

## whether or not to perform log(base2) transformation ##
LOG_TRANSFORM_DNA = false
LOG_TRANSFORM_RNA = false
LOG_TRANSFORM_PROT = false

## parameters within each group ##
MIN_OBS = 10
MIN_PROP = 0.8
### only one of the two has to be specified, MIN_PROP = 0.5 and MIN_OBS=1 by default ###
### if both specified, apply both simultaneously ###

## Apply KNN to impute for missing values ##
KNN_IMPUTE = false
MAX_BLOCKSIZE = 1500

## Specify if there is different directions in the type of interaction edges ##
INTERACT_SIGN = false

ANALYSE_DNA = true
## if true, filename should be provided for DNA_file
## if protein-lvl data is not available, one can also replace protein with rna-level data

## parameter for shrunken centroid ##
CROSS_VALIDATION = false
CV_FOLD = 10
MIN_THRES = 4.0

```

Replace with imputed and standardized files produced from previous run

Turn off options for standardization and log-transformation

Turn off option for imputation to reduce computation time

Re-specify threshold and turn off option for cross-validation

Figure 5: Screen shot of input parameter file after respecifying threshold of 4.0.

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