# MetaboAnalystを用いた PCA, PLS-DA

https://www.metaboanalyst.ca/

## データファイルの作り方

各サンプルが各行になるようにする。行と列を入れ替えることも可能 1列目にID、2列目にサンプルの分類情報、3 列目以降に変数データを記入する

id	class	Q1	Q2	Q3	Q4	Q5	Q6	Q7	Q8
1	M	100	100	50	70	10	100	100	
2	F	80	100	90	100	0	90	100	
3	F	50	100	100	100	0	70	80	
4	F	100	100	50	50	0	70	100	
5	F	94	80	98	30	60	98	99	
6	F	99	100	80	70	0	10	60	
7	M	50	40	80	50	0	80	40	
8	F	55	50	69	100	0	0	15	
9	F	25	0	100	100	0	25	75	
10	F	30	75	80	60	0	90	80	
11	M	85	100	85	90	0	100	40	転置可能
12	M	23	74	87	72	0	68	82	
13	F	70	100	80	100	10	60	60	
14	F	55	90	100	80	30	50	42	
15	F	50	80	100	80	0	30	100	
16	F	90	50	70	70	0	0	40	
17	M	80	30	100	100	2	30	95	

## データファイルの作り方

カンマ区切りファイル(csv)にしたいデータだけを記載した、エクセルシートを作る。

そのシートを閲覧している状態で、 「名前を付けて保存」から、カンマ区切り (csv)を選んでで保存する。

## MetaboAnalystへアクセス

#### click here to startをクリック



MetaboAnalyst - statistical, functional and integrative analysis of metabolomics data

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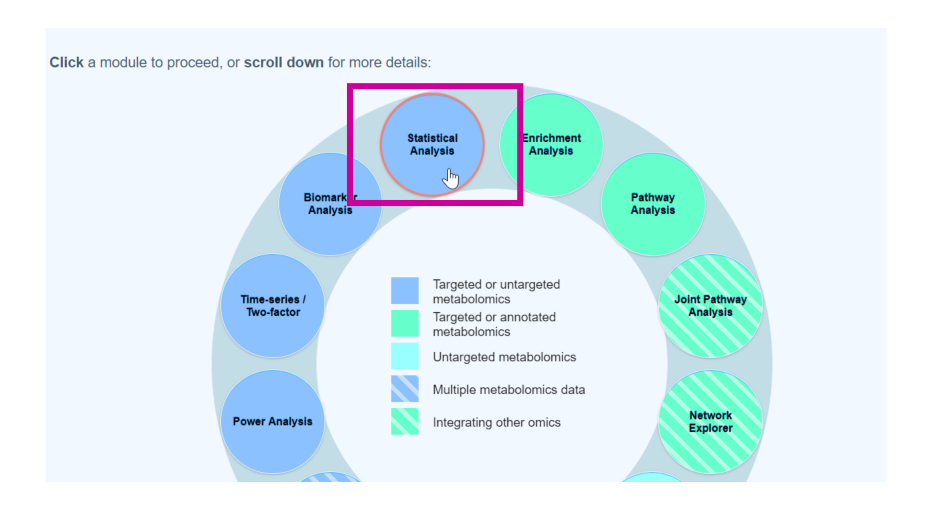


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- · Check out OmicsNet for multi-omics data integration via 3D network visual analytics;
- Check out <u>MicrobiomeAnalyst</u> for comprehensive analysis of microbiome data;
- Check out <u>NetworkAnalyst</u> for comprehensive gene expression profiling & meta-analysis;
- Users can upload MS peak intensity tables and test enriched pathways for any peak cluster of interest (MS Peaks to Pathways) (09/13/2019); 
  \*\*Note: The content of the content of
- Enhanced enrichment network view for metabolite set enrichment analysis (06/25/2019);
- Added mixed mode suppport to allow combining peaks from both postive and negative ion modes (MS peaks to pathways) (06/10/2019);
- Upgraded the web framework (05/27/2019);
- Improved box plot graphics based on user feedback (05/06/2019);
- Enhanced the summary plot for individual metabolite set in Enrichment Analysis module (04/23/2019);
- Enhanced support for various input formats and added interactive result summary for MS Peaks to Pathways module (04/10/2019);
- Added support for synchronized 3D visual exploration for PCA/PLS-DA scores and loading plots (04/07/2019);
- Added support for the GSEA algorithm (MS Peaks to Pathways) (03/22/2019);
- The download table in compound ID conversion now contains SMILES (03/15/2019);
- Enhanced ID mapping for KEGG compounds (03/11/2019);
- Enhanced KEGG pathway generation to address the occasional failure issue during peak time (01/24/2019);
- Fixed the issue for compound view in Pathway Analysis module (01/07/2019);
- Enhanced support for parsing data input and data editing (01/02/2019);

Read more .....

### Statistical Analysisをクリック



### 準備したCSVファイルをアップロード

#### 必要に応じて、Formatなどを選択する

#### Submitボタンを押す

#### 1) Upload your data

Tab-delimited	text (.txt) or comma-separated values (.csv) file:	
Data Type:	Concentrations	
Format:	Samples in rows (unpaired)	Submit
Data File:	ファイルを選択 okonomi.csv	

### データチェックが行われる 問題なければskipで先に進む

#### **Data Integrity Check:**

- 1. Checking the class labels at least three replicates are required in each class.
- 2. If the samples are paired, the pair labels must conform to the specified format.
- 3. The data (except class labels) must not contain non-numeric values.
- 4. The presence of missing values or features with constant values (i.e. all zeros)

#### Data processing information:

Checking data content ...passed

Samples are in rows and features in columns

The uploaded file is in comma separated values (.csv) format.

The uploaded data file contains 33 (samples) by 29 (compounds) data matrix.

Samples are not paired.

2 groups were detected in samples.

Only English letters, numbers, underscore, hyphen and forward slash (/) are allowed.

Other special characters or punctuations (if any) will be stripped off.

All data values are numeric.

A total of 0 (0%) missing values were detected.

By default, these values will be replaced by a small value.

Click Skip button if you accept the default practice

Or click **Missing value imputation** to use other methods

Missing value estimation

Skip

## 必要に応じて、データの標準化(normalization)、 変換(transformation)、スケーリング方法を選択する

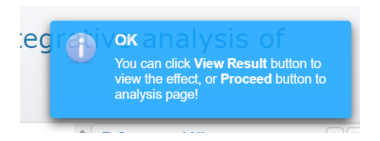
#### Normalizeボタンを押す

#### Normalization overview:

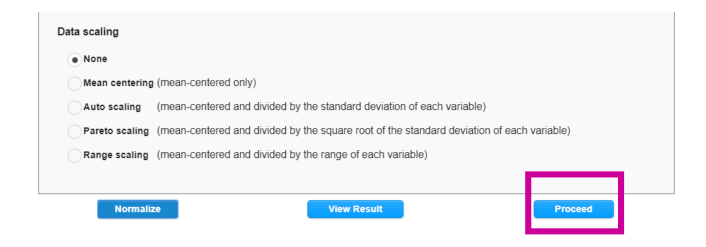
The normalization procedures are grouped into three categories. The sample normalization allows general-purpose adjustment for differences among your sample; data transformation and scaling are two different approaches to make individual features more comparable. You can use one or combine them to achieve better results.

Sample normalization	
None	
Sample-specific normalization (i.e. weight, volume	e) <u>Specify</u>
Normalization by sum	
Normalization by median	
Normalization by reference sample (PQN)	Specify
Normalization by a pooled sample from group	Specify
Normalization by reference feature	Specify
Quantile normalization	
None  Log transformation  Cube root transformation (generalized logarithm transformation (takes the cube root of data.)	
Data scaling	
None	
Mean centering (mean-centered only)	
Auto scaling (mean-centered and divided by the	· ·
	square root of the standard deviation of each variable)
Range scaling (mean-centered and divided by the	range of each variable)
Normalize	Tiew Result Proceed

#### 右上にこのウィンドウが出ればOK



#### アクティブになったProceedボタンを押す



### 主成分分析 Principal Component Analysis (PCA) をクリック

#### Select an analysis path to explore:

#### **Univariate Analysis**

Fold Change Analysis T-tests Volcano plot

One-way Analysis of Variance (ANOVA)

Correlation Analysis Pattern Searching

#### **Chemometrics Analysis**

Principal Component Analysis (PCA)

Partial Least Squares - Discriminant Analysis (PLS-DA)

Sparse Partial Least Squares - Discriminant Analysis (sPLS-DA)

Orthogonal Partial Least Squares - Discriminant Analysis (orthoPLS-DA)

#### **Feature Identification**

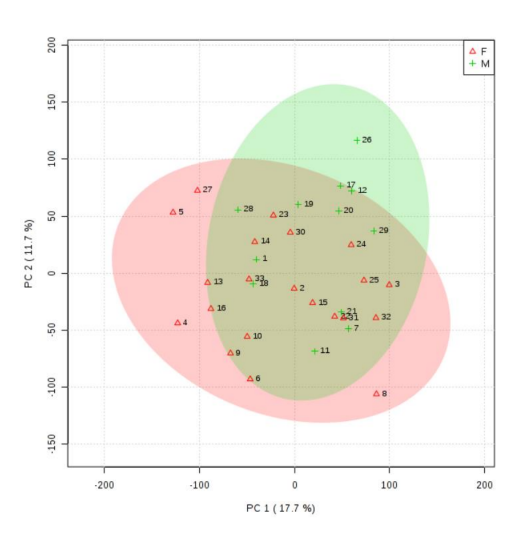
Significance Analysis of Microarray (and Metabolites) (SAM)

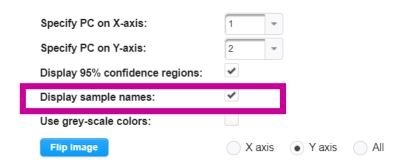
Empirical Bayesian Analysis of Microarray (and Metabolites) (EBAM)

#### 結果画面が表示される



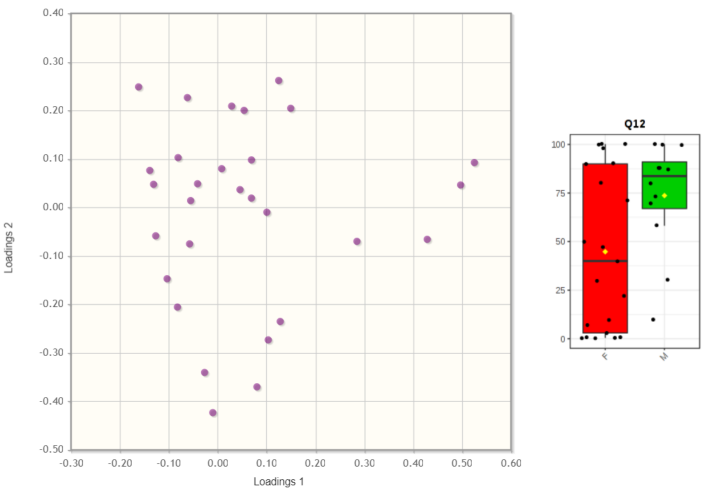
#### **2D Scores Plot**





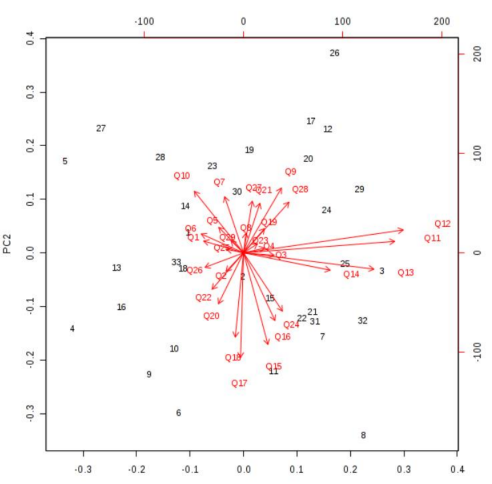
Display sample nameにチェックが入っていると、サンプル名が一緒に表示される

### **Loadings Plot**



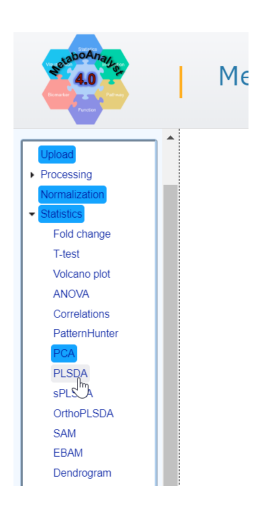
プロットをクリックすると、 その変数の特徴が表示される。

#### **Biplot**

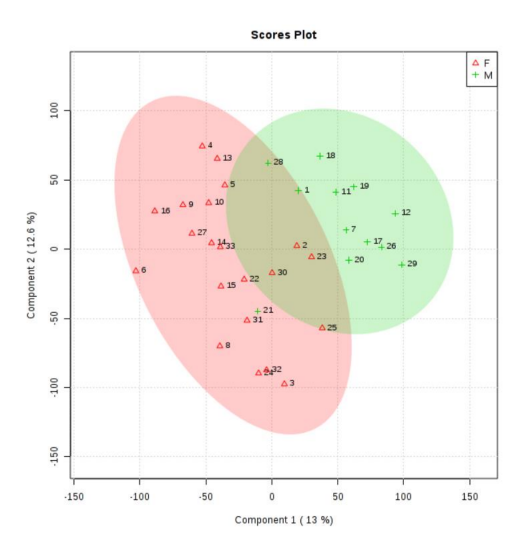


スコアプロットとローディングプロットが一緒に描かれたもの。

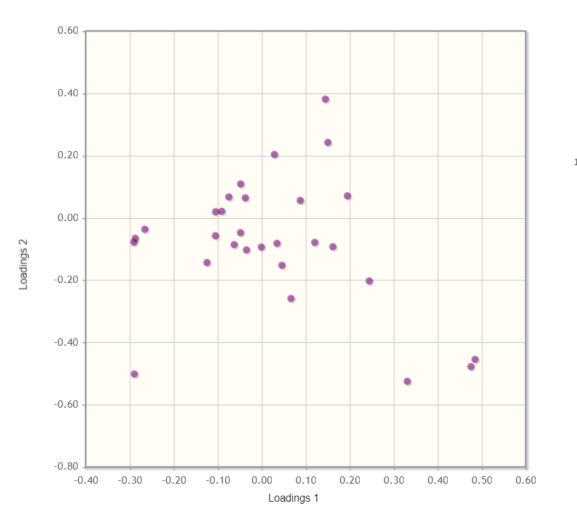
#### **PLSDA**

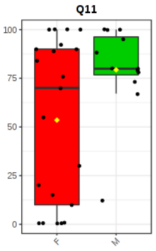


#### **2D Scores Plot**

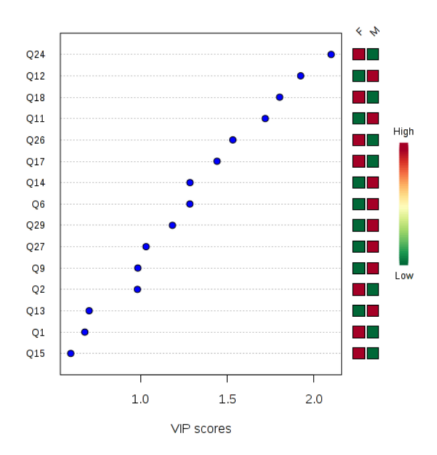


### **Loadings Plot**





### VIP値



### 累積寄与率と構築したモデルの精度(Q)

