

Energy Metabolism Targeted Metabolomics Assay Final Report

Metware Biotechnology Inc.

www.metwarebio.com



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MWY-23-XXX Energy Targeted Metabolomics Assay Final Report

1 Abstract

Changes in physiological activity can change the metabolite profile of an organism. Targeted quantitative detection technology allows sensitive qualitative annotation and highly accurate quantitative analysis of a set of metabolites. MetwareBio has established an LC-MS/MS based analytical method that can quantify 68 energy related metabolites.

For this project,24 samples were divided into 4 groups. A total of 53 metabolites were detected based on UPLC-MS/MS system.

2 The experimental process

Liquid chromatography-tandem mass spectrometry (LC-MS/MS) can detect and quantify compounds with high polarity and poor thermal stability, and accurately quantify them. The overall process is as follows:



Fig 1: Flow chart of metabolomics analysis

Compounds to be detected:

Table 1:	List of	compounds	in	the	panel	
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Number	Compounds	Index
1	Pyruvic-acid	Pyruvic-acid
2	Serine	Serine
3	L-Glutamic-acid	L-Glutamic-acid
4	Threonine	Threonine
5	Lysine	Lysine
6	Tyrosine	Tyrosine



Number	Compounds	Index
7	Arginine	Arginine
8	Ornithine	Ornithine
9	L-Leucine	L-Leucine
10	Glutamine	Glutamine
11	L-Alanine	L-Alanine
12	Succinic Acid	Succinic-Acid
13	Alpha-Ketoglutaric-Acid	Alpha-Ketoglutaric-Acid
14	L-Asparagine	L-Asparagine
15	Adenine	Adenine
16	Inosine	Inosine
17	3-phenyllactic-acid	3-phenyllactic-acid
18	Citric-acid	Citric-acid
19	Lactate	Lactate
20	ADP	ADP
21	Fumaric-acid	Fumaric-acid
22	Uracil	Uracil
23	Guanosine	Guanosine
24	c-di-AMP	c-di-AMP
25	D-Glucose-6-phosphate	D-Glucose-6-phosphate
26	Cyclic-AMP	Cyclic-AMP
27	Fructose-1,6-bisphosphate	Fructose-1,6-bisphosphate
28	Glycerol-3-phosphate	Glycerol-3-phosphate
29	Phosphoenolpyruvic-acid	Phosphoenolpyruvic-acid
30	6-Phosphogluconic-acid	6-Phosphogluconic-acid
31	D-Erythrose 4-phosphate	D-Erythrose-4-phosphate
32	Dihydroxyacetone-phosphate	Dihydroxyacetone-phosphate
33	Isocitric-acid	Isocitric-acid
34	Flavin-mononucleotide	Flavin-mononucleotide
35	AMP	AMP
36	dCMP	dCMP
37	dAMP	dAMP
38	IMP	IMP
39	UMP	UMP
40	dTMP	dTMP
41	Trehalose-6-phosphate	Trehalose-6-phosphate
42	Guanosine-diphosphate	Guanosine-diphosphate
43	L-Cystine	L-Cystine
44	Acetyl-CoA	Acetyl-CoA
45	Argininosuccinic-acid	Argininosuccinic-acid
46	Phosphorylethanolamine	Phosphorylethanolamine
47	L-citrulline	L-citrulline
48	ATP	ATP
49	Oxaloacetate	Oxaloacetate
50	D-Glucose-1-phosphate	D-Glucose-1-phosphate
51	Sedoheptulose-7-phosphate	Sedoheptulose-7-phosphate
52	D-Ribulose-5-phosphate	D-Ribulose-5-phosphate
53	dUMP	dUMP

Table 1: List of compounds in the panel



Number	Compounds	Index
54	UDP-GlcNAc	UDP-GlcNAc
55	BPG	BPG
56	2-Phospho-D-glyceric acid	2-Phospho-D-glycerate
57	Succinyl-CoA	Succinyl-CoA
58	cis-Aconitic-acid	cis-Aconitic-acid
59	Itaconic-acid	Itaconic-acid
60	Malic-acid	Malic-acid
61	D-Fructose-6-phosphate	D-Fructose-6-phosphate
62	Glyceraldehyde-3-phosphate	Glyceraldehyde-3-phosphate
63	NicotinaMide-adenine-dinucleotide(NAD)	NicotinaMide-adenine-dinucleotide(NAD)
64	Dihydronicotinamide-adenine-dinucleotide-	Dihydronicotinamide-adenine-dinucleotide-
	phosphate(NADPH)	phosphate(NADPH)
65	D(+)-Glucose	D(+)-Glucose
66	L-Aspartate	L-Aspartate
67	Xyluose-5-phosphate	Xyluose-5-phosphate
68	3-phosphoglycerate	3-phosphoglycerate

Table 1: List of compounds in the panel

Original file path: Final report/data/metware_component.xlsx

2.1 Sample information

This project has 24 samples divided into 4 groups. Sample information is shown in the following table:

Species	Tissues	MW_ID	Sample_ID
_	_	A1	Al
_	_	A2	A2
_	_	A3	A3
_	_	A4	A4
		A5	A5
_	_	A6	A6
-	-	B1	B1
_	_	B2	B2
_	_	B3	B3
_	_	B4	B4
_	_	B5	B5
_	_	B6	B6
-	-	Cl	C1
-	-	C2	C2
-	-	C3	C3
-	-	C4	C4
-	-	C4	C4 C5
_	_		
-	-		
-	-		
_	_	D2	D2

Table 2: Sample information table



Species	Tissues	MW_ID	Sample_ID
-	-	D3	D3
-	-	D4 D5	D4 D5
-	-	D6	D6

Table 2: Sample information table

Original file path: Final report/0.data/sample_info.xlsx

2.2 Reagents and instruments

Instrument	Model	Manufacturer
LC-MS/MS	Triple Quad 6500+	SCIEX
Centrifuge	5424R	Eppendorf
Electronic balance	AS 60/220.R2	RADWAG
Ball mill instrument	MM400	Retsch
Multitube vortex oscillator	MIX-200	ShangHaiJingXin
Ultrasonic cleaning apparatus	CD-F15	Olenyer

Table 3: Instrument information

Table 4:	Information	of standards	and reagents
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Reagent	level	Manufacturer
Methanol	HPLC	Thermo fisher
Acetonitrile	HPLC	Thermo fisher
Formic acid	HPLC	Thermo fisher
Ammonia	HPLC	Thermo fisher
Ammonium acetate	LC-MS	Sigma-Aldrich
Chemical standard	99%	Sigma-Aldrich/Zhenzhun.etc

2.3 Sample extraction process

Approximately 0.05 g of the thawed ground sample was mixed with 500 μ L of 70% methanol/water. The sample was vortexed for 3 min at 2500 r/min and centrifuged at 12000 rpm for 10 min at 4°C. About 300 μ L of the supernatant was transferred into a new centrifuge tube and placed in -20°C for 30 min. The supernatant was centrifuged again at 12000 rpm for 10 min at 4°C. About 200 μ L of the supernatant were transferred through Protein Precipitation Plate for further LC-MS analysis.

2.4 Chromatography-mass spectrometry acquisition conditions

The sample extracts were analyzed using an LC-ESI-MS/MS system (Waters ACQUITY H-Class, https://www.waters.com/nextgen/us/en.html; MS, QTRAP® 6500+ System, https://sciex.com /). The analytical conditions were as follows.

UPLC Conditions:

Amide method: HPLC: column, ACQUITY UPLC BEH Amide (i.d.2.1×100 mm, 1.7 μ m); solvent system, water with 10mM Ammonium acetate and 0.3% Ammonium hydroxide (A), 90% acetonitrile/water (V/V)(B); The gradient was started at 95% B (0-1.2 min), decreased to 70% B (8 min),50% B (9-11 min), finally ramped back to 95% B (11.1-15 min); flow rate, 0.4 mL/min; temperature, 40°C; injection volume: 2 μ L.

ESI-MS/MS Conditions:

Linear ion trap (LIT) and triple quadrupole (QQQ) scans were acquired on a triple quadrupole-linear ion trap mass spectrometer (QTRAP), QTRAP® 6500+ LC-MS/MS System, equipped with an ESI Turbo Ion-Spray interface, operating in both positive and negative ion mode and controlled by Analyst 1.6.3 software (Sciex). The ESI source operation parameters were as follows: ion source, ESI+/-; source temperature 550 °C; ion spray voltage (IS) 5500 V (Positive), -4500 V (Negative); curtain gas (CUR) was set at 35 psi, respectively. Metabolites were analyzed using scheduled multiple reaction monitoring (MRM). Data acquisitions were performed using Analyst 1.6.3 software (Sciex). Multiquant 3.0.3 software (Sciex) was used to quantify all metabolites. Mass spectrometer parameters including the declustering potentials (DP) and collision energies (CE) for individual MRM transitions were done with further DP and CE optimization. A specific set of MRM transitions were monitored for each period according to the metabolites eluted within this period.

2.5 Qualitative and quantitative principles of metabolites

Metabolites were quantified by multiple reaction monitoring (MRM) using triple quadrupole mass spectrometry. In MRM mode, the first quadrupole screened the precursor ions for the target substance and excluded ions of other molecular weights. After ionization induced by the impact chamber, the precursor ions were fragmented, and a characteristic fragment ion was selected through the third quadrupole to exclude the interference of non-target ions. After obtaining the metabolite spectrum data from different samples, the peak area was calculated on the mass spectrum peaks of all substances and analyzed by standard curves.





Fig 2: Schematic diagram of multiple reaction monitoring mode by mass spectrometry

3 Data evaluation

3.1 Data pre-processing

Analyst 1.6.3 was used to process mass spectrum data. The following figure shows the total ions current (TIC) and MRM metabolite detection multi-peak diagram (XIC) of the mixed QC samples. The X-axis shows the Retention time (RT) from metabolite detection, and the Y-axis shows the ion flow intensity from ion detection (intensity unit: CPS, count per second).



Fig 3: Total ion current diagram of mixed phase mass spectrum analysis

Original file path: Final report/0.data/QC/*QC_MS_TIC.png





Fig 4: Extraction ion flow chromatogram

Original file path: Final report/0.data/QC/*MRM_detection_of_multimodal_maps*

The mass spectrometry data was analyzed using MultiQuant 3.0.3 software. The mass spectrum peaks detected in different samples were scored and corrected based on retention time and peak shape of the standard. The figure below shows the correction results of quantitative analysis of a substance randomly selected from different samples.



Fig 5: Scoring correction diagram for quantitative analysis of metabolites Note: The figure shows the quantitative analysis integral correction results of randomly selected metabolites in different samples. The x-axis is the retention time (min) of metabolite detection, the y-axis is the ion flow intensity (CPS) of a certain metabolite ion detection, and the peak area represents the relative content of the substance in the sample.

Original file path: Final report/0.data/QC/*Integral_correction.png

3.2 Standard Solution Preparation

Standards were prepared at 0.01 ng/mL, 0.02 ng/mL, 0.05 ng/mL, 0.1 ng/mL, 0.2 ng/mL, 0.5 ng/mL, 1 ng/mL, 2 ng/mL, 5 ng/mL, 10 ng/mL, 20 ng/mL, 50 ng/mL, 100 ng/mL, 200 ng/mL, 500 ng/mL, 1000 ng/mL, 2000 ng/mL, 5000 ng/mL, and 10000 ng/mL. NADPH was prepared at 0.1 ng/mL, 0.2 ng/mL, 0.5 ng/mL, 1 ng/mL, 2 ng/mL, 5 ng/mL, 10 ng/mL, 20 ng/mL, 50 ng/mL, 100 ng/mL, 2000 ng/mL, 500 ng/mL, 1000 ng/mL, 2000 ng/mL, 500 ng/mL, 500 ng/mL, 1000 ng/mL, 2000 ng/mL, 5000 ng/mL, 500 ng/mL, 1000 ng/mL, 1000 ng/mL, 2000 ng/mL, 5000 ng/mL, 5000 ng/mL, 500 ng/mL, 1000 ng/mL, 1000 ng/mL, 50000 ng/mL, 50000 ng/mL, 5000 ng/mL. Mass spectral peak intensity data were collected at each concentration to generate the calibration curve. The standard curves of different substances were plotted with the external standard concentration as the horizontal coordinate and the peak area of the external standard as the vertical coordinate. The equation of calibration curve are shown in the following table:

Index	Class	RT	Equation
L-Aspartate	Amino Acid metabolomics	5.36	y = 6319.39470 x - 1.95245e5
Glutamine	Amino Acid metabolomics	5.75	y = 22513.29896 x - 19002.34080
Ornithine	Amino acids	10.11	y = 3.13681e4 x + 4.62215e4
Arginine	Amino acids	10.61	y = 1.57487e5 x + 9.68610e4
Lysine	Amino acids	10.77	y = 5.80830e4 x - 7.67596e4
cis-Aconitic-acid	Amino acids	2.54	y = 1.05156e5 x - 1.35894e6
Itaconic-acid	Amino acids	2.54	y = 3.80208e4 x - 3.87019e5
L-Leucine	Amino acids	2.68	y = 5.61828e4 x - 6.89072e4
Tyrosine	Amino acids	3.80	y = 5829.81773 x - 5.74400e5
Malic-acid	Amino acids	4.32	y = 8.91447e4 x - 7.85966e6

Table 5: Equation of calibration curve

Final report/0.data/equation.xlsx

3.3 Quantification Results

Concentrations of each compound was obtained by substituting integrated peak area of all the detected samples into the equation of calibration curve .

Concentration of solid sample (ng/g) = c*V/1000/m

c: the concentration obtained by substituting the sample peak area into the equation of calibration curve (ng/mL);

V: the volume of extraction solution (μ L);

m: the mass of the sample (g).

The metabolite ID, concentration and corresponding metabolite names of some metabolites detected in this experiment are shown in the following table:



Index	A1	A2
L-Aspartate	0.07732	0.07208
Glutamine	0.00503	0.00537
Ornithine	0.05830	0.05281
Arginine	0.07301	0.08033
Tyrosine	0.05797	0.05959
Lysine	0.00653	0.00690
Threonine	0.06202	0.06248
L-Glutamic-acid	0.06404	0.06089
Serine	0.10439	0.09678
L-Alanine	0.07713	0.07291

Table 6: Statistical Table of metabolite quantity

Original file path: Final report/0.data/*level.xlsx

3.4 Sample Quality Control Analysis

3.4.1 Total Ion Chromatogram Analysis

Using the mixed solution as the QC sample, one QC sample was inserted every 10 detection samples for analysis during the detection by the system. The stability of the device during the detection of the project can be assessed by analyzing the overlapped total ion flow chromatograms (TICs) obtained from the mass spectrometry detection and analysis of the same QC samples. The high stability of the testing device is a vital safeguard for the reproducibility and reliability of the data.



Fig 6: TIC overlap diagram detected by QC sample essence spectrum Note: Superimposed spectrum from different QC samples. The results showed that the spectrum of total ion flow were highly consistent indicating that the signal stability was good when the same sample was detected at different times by mass spectrometry.

Original file path: Final report/0.data/picture/*QC_MS_tic_overlap*

3.4.2 QC Sample correlation assessment

Pearson correlation analysis was performed on the QC samples. The closer the |r| to 1, the higher the correlation between two samples. The correlation results can be seen in the figure below.





Note: Diagonal squares represent QC samples name; Left diagonal box represent scatter diagram of QC samples . Both x-axis and y-axis represent metabolite content. Each dot in the diagram represents a metabolite. Right diagonal box represents correlation coefficients of QC samples .

Original file path/1.Data_Assess/pcc/*mix*

3.4.3 CV value distribution of all samples

The Coefficient of Variation (CV) value is the ratio between the standard deviation of the original data and the mean of the original data, which can reflect the degree of data dispersion. The Empirical Cumulative Distribution Function (ECDF) can be used to analyze the frequency of CV of substances that is smaller than the reference value. The higher the proportion of substances with low CV value in QC samples is, the more stable the experimental data is. The proportion of substances with CV value less than 0.3 in QC samples was higher than 80%, indicating that the experimental data were relatively stable. The proportion of substances with CV value less than 0.2 in QC samples was higher than 80%, indicating that the experimental data were very stable.





Fig 8: CV distribution of each group

Note: The X-axis represents the CV value, the Y-axis represents the proportion of metabolites with CV value less than a corresponding reference value. Different colors represent different sample groups. QC indicates quality control samples. The two dash lines on X-axis correspond to 0.2 and 0.3; the two dash lines on Y-axis correspond to 80%.

Original file path: Final report/1.Data Assess/CV/*ECDF*

3.5 Principal Component Analysis (PCA)

3.5.1 Principles of principal component analysis

Multivariate statistical analysis can simplify complex high-dimensional data while preserving the original information to the maximum extent by establishing a reliable mathematical model to summarize the characteristics of the metabolic spectrum. Among them, Principal Component Analysis (PCA) is an unsupervised pattern recognition method for statistical analysis of multidimensional data. Through orthogonal transformation, a group of variables that may be correlated are converted into a group of linear unrelated variables that are called principal components. This method is used to study how a few principal components may reveal the internal structure of multiple variables, while keeping the original variable information (Eriksson et al., 2006). The first principal component (PC1) represents the most variable features in the multidimensional data matrix, PC2 represents the second most variable feature in the data, and so on. prcomp function of R software (www.r-project.org/) was used with parameter scale=True indicating unit variance Scaling (UV) for normalizing the data. See appendix for details of PCA calculation.

3.5.2 Principal component analysis of the sample population

Principal component analysis (PCA) was performed on all the samples (including QC samples) to examine the overall differences between each group and the variation between samples within a group. QC is the Quality control sample mentioned above. PCA plot for the first two principal components is as follows:







diagram of quality spectrum data of each group of samples and quality control sample Note: PC1 represents the first principal component and PC2 represents the second principal component. Percentage represents the interpretation rate of the principal component to the data set. Each dot in the figure represents a sample, and samples in the same group are indicated in the same color.

Original file path : Final report /1.Data_Assess/*all_pca*

3.5.3 Principal component univariate statistical process control

We plotted the sample control diagram based on principle component analysis results. Each point in the control chart represents a sample, and the X-axis is the injection order of the sample. Due to changes in the instrument, the points on the chart may fluctuate up and down. Generally, PC1 of the QC sample should be within 3 standard deviations (SD) from the normal range.





Fig 10: PC1 control diagram of population sample

Note: In the figure, the X-axis is the injection order of the sample, and the Y-axis reflects the PC1 value. The yellow and red lines define plus or minus 2 and 3 standard deviations respectively. The green dots represent QC samples and the black dots represent test samples.

Original file path: Final report/1.Data Assess/pca/*PC1 QCC*

3.6 Hierarchical Cluster Analysis

3.6.1 Principles of cluster analysis

Hierarchical Cluster Analysis (HCA) is a type of multivariate statistical analysis method. The samples are classified according to their features such that highest homogeneity is achieved between sample from the same group and highest heterogeneity is achieved between samples from different groups. In this report, the compound quantification data was normalized (Unit Variance Scaling, UV Scaling) and heatmaps were drawn by R software Pheatmap package. Hierarchical Cluster Analysis (HCA) was used to cluster the samples.







Note: X-axis indicates the sample name and the Y-axis are the metabolites. Group indicates sample groups. Z-Score indicates the relative quantification of each metabolite with red representing higher content and green representing lower content. Cluster analysis was performed on both metabolites (vertical cluster tree) and samples (horizontal cluster tree). "all_heatmap_class" : Heat map based on metabolite classification; "all_heatmap_no_cluster" : Showing only heatmap.

Original file path: Final report /1.Data_Assess/*all_heatmap*

4 Analysis results

4.1 Principal component analysis of sample groups

4.1.1 Principal component analysis between sample groups

Principal component analysis was first performed on each pair of sample groups to examine the degree of variation between different groups and between samples within the group.





Fig 12: Principal component analysis of different groups

Note: Each group has a PCA plot, PC1 represents the first principal component, PC2 represents the second principal component, and the percentages on the axis represents the interpretation rate of the principal component to the data set. Each dot in the figure represents a sample, samples in the same Group are represented by the same color, and Group is a grouping.

The three-dimensional PCA result is shown in the figure below:



Fig 13: Three-dimensional PCA plot of different groups Note: PC1 represents the first principal component, PC2 represents the second principal component, and PC3 represents the third principal component.

The explainable variation of the first five principal components is shown in the figure.





Fig 14: The explainable variation of the first five principal components Note: The X-axis represents each principal component, the Y-axis represents the explainable variation, the left figure represents the cumulative explainable variation, and the right figure represents the explainable variation of each principal component

Principal component analysis of different groups:Original file path: Final report/2.Basic_analysis/Difference_analysi ID*_vs_group-ID*/pca/group-ID*_vs_group-ID*_pca.*;

Three-dimensional PCA plot of different groups:Original file path: Final report/2.Basic_analysis/Difference_analysis ID*_vs_group-ID*/pca/group-ID*_vs_group-ID*_pca3D.*

The explainable variation of the first five principal components:Original file path: Final report/2.Basic_analysis/Difference_analysis/group-ID*_vs_group-ID*/pca/group-ID*_vs_group-ID*_pcaVar.*

4.2 Discriminant Analysis by Orthogonal Partial Least Squares (OPLS-DA)

PCA analysis is often insensitive to variables with small correlation. In contrast, partial least squaresdiscriminant analysis (PLS-DA) is a multivariate statistical analysis method with supervised pattern recognition, in which components in independent variable X and dependent variable Y are extracted to calculate the correlation between components. Compared with PCA, PLS-DA can maximize the difference between groups and facilitate the search for differential compounds. Orthogonal partial least squares discriminant analysis (OPLS-DA) combines orthogonal signal correction (OSC) and PLS-DA method, which can decompose the x-matrix information into two types (1. information related to Y and 2. irrelevant information) and filter the differential variables by removing the irrelevant differences.

The OPLSR.Anal function in the R package MetaboAnalystR was used for this analysis. The following table shows a partial result from the OPLS-DA model:



Index	Compounds	Туре
L-Aspartate	L-Aspartate	up
Glutamine	Glutamine	down
Ornithine	Ornithine	insig
Arginine	Arginine	insig
Tyrosine	Tyrosine	up
Lysine	Lysine	down
Threonine	Threonine	up
L-Glutamic-acid	L-Glutamic-acid	insig
Serine	Serine	up
L-Alanine	L-Alanine	insig

Table 7: Partial results of OPLS-DA

Original file path: The calculation results of all metabolites of OPLS-DA were compared in groups: /2.Basic_analysis/Difference_analysis/group-ID*_vs_group-ID*/group-ID*_vs_group-ID*_info.xlsx.

4.2.1 Principles of OPLS-DA model

During OPLS-DA modeling, the X matrix information is decomposed into information related to Y and information unrelated to Y. Among them, the variable information related to Y is the predicted principal component, and the information unrelated to Y is the orthogonal principal component (Thevenot et al., 2015).



Fig 15: OPLS-DA score diagram

Note: The X-axis represents the predicted principal component, and the difference between groups can be seen in the horizontal direction. The Y-axis represents the orthogonal principal component, and the vertical direction shows the difference within the group. Percentage indicates the degree to which the component explains the data set. Each dot in the figure represents a sample, samples in the same Group are represented by the same color, and Group indicates sample groups.

Original file path:Final report/2.Basic_analysis/Difference_analysis/group-ID*_vs_group-ID*/opls/group-ID* vs_group-ID* opls_score.*.

4.2.2 OPLS-DA model validation

The prediction parameters of the evaluation model are R^2X , R^2Y and Q^2 , where R^2X and R^2Y represent the explanatory rate of the model to X and Y matrix respectively, and Q^2 represents the predictability of the model. The closer these three indicators are to 1, the more stable and reliable the model is. $Q^2 > 0.5$ can be considered as an effective model, and $Q^2 > 0.9$ can be considered as an excellent model. The following figure shows the OPLS-DA validation plot with the horizontal coY-axis indicating the model R^2Y , Q^2 values, and the vertical coY-axis is the frequency of the model classification effect. The model performs bootstrapping 200 times and if Q^2 's P = 0.02, it indicates that the prediction ability of four random grouping models is better than that of the OPLS-DA model in the Permutation detection. If R^2Y 's P = 0.545, it indicated that there were 109 random grouping models in the Permutation detection, whose explanation rate of Y matrix was better than that of the OPLS-DA model. In general, P < 0.05 is the best model.



Fig 16: OPLS-DA verification diagram

Original file path:Final report/2.Basic_analysis/Difference_analysis/group-ID*_vs_group-ID*/opls/group-ID*_vs_group-ID*_opls_permutation.*.

4.2.3 OPLS-DA S-plot

The figure below shows the OPLS-DA S-plot. The horizontal axis is the covariance between the principal components and metabolites, the vertical axis indicates the correlation coefficient between the principal components and the metabolites. The closer the points are to the top right corner or bottom left corner, the more significant the difference in metabolite abundance. Red dots indicate metabolites with VIP value > 1 and green dots indicate metabolites with VIP value <= 1.







Original file path:Final report/2.Basic_analysis/Difference_analysis/group-ID*_vs_group-ID*/opls/group-ID*_vs_group-ID*_opls_splot.*.

4.2.4 Dynamic distribution of metabolite content differences

To show the overall compound abundance distribution in the samples, compounds were sorted and plotted based on fold-change values from small to large. The distribution of the ranked compounds is shown below with the top 10 up-regulated and top 10 down-regulated compound labelled.



Fig 18: Dynamic distribution of metabolite content differences Note: In the figure, the X-axis represents the rank number of metabolites based on FC value. The Y-axis represents the log_2FC value. Each point represents a metabolite. The green points represent the top 10 down-regulated metabolites and the red points represent the top 10 up regulated metabolites.

Final report/2.Basic_analysis/Difference_analysis/group-ID*_vs_group-ID*/distribution/group-ID*_vs_group-ID*fc_distribution*

4.3 Differential metabolite screening

It is often necessary to combine univariate statistical analysis and multivariate statistical analysis for large high dimensional datasets such as metabolomics datasets to accurately identify differential metabolites. Univariate statistical analysis methods include parametric test and nonparametric test. Multivariate statistical analysis methods include principal component analysis and partial least square discriminant analysis. Based on the results of OPLS-DA (biological repetition \geq 2), multivariate analysis of Variable Importance in Projection (VIP) from OPLS-DA modeling was used to preliminarily select differential metabolites from different samples. The fold-change and statistical significance (p-value) from univariate analysis can be used in conjunction to further identify differential metabolites. If biological replicates were < 3, differential metabolites are screened based on Fold Change value. If there were \geq 3 biological replicates, VIP and P-values were used in combination to screen for differential metabolites. The detailed screening criteria is as follows:

For two sets of comparisons:

1.Metabolites with VIP > 1 were selected. VIP value represents the effect of the differences between groups for a particular metabolite in various models and sample groups. It is generally considered that the metabolites with VIP > 1 have significant difference.

2.Metabolites with Fold Change \geq 2 and Fold Change \leq 0.5 were considered as significant and selected.

A partial result from the screening criteria is seen below:

Index	Compounds	Туре
L-Aspartate	L-Aspartate	up
Glutamine	Glutamine	down
Tyrosine	Tyrosine	up
Lysine	Lysine	down
Threonine	Threonine	up
Serine	Serine	up
Itaconic-acid	Itaconic-acid	down
Malic-acid	Malic-acid	down
L-citrulline	L-citrulline	down
Dihydronicotinamide-adenine-dinucleotide- phosphate(NADPH)	Dihydronicotinamide-adenine-dinucleotide- phosphate(NADPH)	up

Table 8: Screening results of differential metabolites

Final report/2.Basic_analysis/Difference_analysis/group-ID*_vs_group-ID*/group-ID*/group-ID*_to_sgroup-ID*filter.xlsx.

4.3.1 Bar chart of differential metabolites

The following figure shows the result of top differentially expressed metabolites in each comparison with fold-change value shown as \log_2 values .





refers to metabolites. Red bars represent up-regulated differential metabolites and green bars represent down-regulated differential metabolites.

Final report/2.Basic_analysis/Difference_analysis/group-ID*_vs_group-ID*/TopFcMetabolites/group-ID*_vs_group-ID*_TopFcMetabolites.*

4.3.2 Differential metabolite radar map

The top 10 differential metabolites based on Fold-change were selected and plotted on the radar plot.





Fig 20: Differential metabolite radar map Note: The grid lines correspond to the log_2FC. The green colored area is formed from the lines connecting the dots

Final report/2.Basic Analysis/Difference analysis/group-ID* vs group-ID*/radarchart/*radarchart**

4.3.3 VIP values of differential metabolites

The top 50 metabolites with the largest VIP value in the OPLS-DA model were selected and plotted.





/2.Basic_analysis/Difference_analysis/group-ID*_vs_group-ID*/vipscore/group-ID*_vs_group-ID*_vipScore.*.

4.3.4 Volcanic plot of differential metabolites

Volcano Plot is mainly used to show the relative differences and the statistical significance of compounds between two groups. We provided the volcano plot of differential compounds using different selection criteria for your consideration. The details of different selection criteria are described in the README document under the volcano plot directory. In addition, the attached results also provided an interactive web version of the volcano plot where you can examine the details of each compound.



Fig 22: Volcanic plot of differential metabolites

Note: Each point in the volcano plot represents a metabolite with green dots represents the down-regulated differential metabolite, red dots represents the up-regulated differential metabolite, and gray dots represents the detected metabolite but show no insignificant difference. The X-axis represents the (log_2FC) value of metabolite between two groups. The further away from 0 on the X-axis, the greater the fold-change between two groups. If the metabolite were screened using VIP + FC + P-value, the Y-axis will represent the level of significant difference (-log_10p-value). The size of each dot represents the VIP value

Final report/2.Basic Analysis/Difference analysis/group-ID* vs group-ID*/vol/*vol *

4.3.5 Heatmap of differential metabolites

In order to observe the fold-change of differential compounds more intuitively, we normalized the abundances using unit variance scaling (UV scaling, see appendix for details of calculation formula) and plotted on a heatmap using pheatmap in R.







Note: The X-axis shows the name of the samples and the Y-axis shows the differential metabolites. Different colors in the heatmap represent the values obtained after normalization and reflect the level of relative quantification. The darker the red, the higher the quantification. In contrast, the darker the green, the lower the quantification. The colored bar on top depicts sample groups. If hierarchical clustering is performed, the clustering tree will be shown on the left. If classification was performed on the metabolites, a colored bar will be shown on the left to depict Level 1 classifications.

Heatmap of differential metabolites:Final report/2.Basic_analysis/Difference_analysis/group-ID*_vs_group-ID*/heatmap/group-ID* vs_group-ID* heatmap.*;

4.3.6 Z-value map of differential metabolites

Z-score plot is to normalize the differential metabolites in different samples by calculating the Z-value. The a-axis represents the z-value, the y-axis represents the differential metabolites, and the dots in different colors represent samples of different groups. The distribution of each differential metabolite among different groups can be seen intuitively. The formula is: $z = (x - \mu) / \sigma$; Where x is a specific score, μ is the mean, and σ is the standard deviation.





Fig 24: Z-value map of differential metabolites

Note: the X-axis is the value of substance content after normalized treatment, the Yaxis is the number of metabolites, and the points in different colors represent different groups of samples.

/2.Basic_analysis/Difference_analysis/group-ID*_vs_group-ID*/zScore/group-ID*_vs_group-ID*_zScore.*.

4.3.7 Correlation analysis of differential metabolites

Compounds may act synergistically or in mutually exclusive relationships amongst each other. Correlation analysis can help measure the compound proximities of significantly different compounds. This analysis will help further understand the mutual regulatory relationship between compounds in the biological process. Pearson correlation was used to perform correlation analysis on the differential compounds identified based on the screening criteria described previously.







Note: The ID of the metabolites are shown on both horizontal and vertical axis. The colors represent the Pearson correlation coefficient (r) with the scale seen on the right (The darker the red, the stronger the positive correlation; the darker the green the stronger the negative correlation). If there are more than 50 differential metabolites, the figure will only show the top 50 metabolites based on VIP values.

Differential metabolite correlation heat map: Final report/2.Basic_analysis/Difference_analysis/group-ID* vs group-ID*/cpdCorr/group-ID* vs group-ID* raw cpdCorr *.*;



Fig 26: Chord diagram of differential metabolites

Note: The outermost layer shows the metabolite ID. The second layer shows log_2FC value, The larger the dot, the larger the log_2FC value; The color for the first and second layer represent Level 1 metabolite classification. The chords in the inner most layer reflect the Pearson correlation between the connected metabolites. Red chords represent positive correlation, and the blue chords represent negative correlation. Only metabolites with $|\mathbf{r}| \ge 0.8$ and p < 0.05 are plotted.



Final report//2.Basic_analysis/Difference_analysis/group-ID*_vs_group-ID*/cpdCorr/group-ID*_vs_group-ID* cpdCorrCir *.*;



Fig 27: Correlation network diagram of differential metabolites Note: The points in the figure represent the various differential metabolites, and the size of the points is related to the Degree of connection. The larger the point, the greater the Degree of connection, i.e. the more points (neighbors) connected to it. Red lines represent positive correlations and blue lines represent negative correlations. Line thickness represents the absolute value of Pearson correlation coefficient. The larger the $|\mathbf{r}|$, the thicker the line. Only metabolites with $|\mathbf{r}| \ge 0.8$ and p < 0.05are plotted.

Final report/2.Basic_Analysis/Difference_analysis/group-ID*_vs_group-ID*/cpdCorr/*network*

4.3.8 Violin plot of differential metabolites

Violin plot is used to display data distribution and its probability density. The box in the middle represents the interquartile range, and the middle box represents the 95% confidence interval. The black horizontal line is the median, and the outer shape represents the distribution density of the data. The following figure shows the result of top 50 differentially compounds with the largest Log_2FC value.



Fig 28: Violin plot of differential metabolites Note: X-axis refers to sample, the Y-axis refers to content.

Final report/2.Basic_analysis/Difference_analysis/group-ID*_vs_group-ID*/fullViolin/group-ID*_vs_group-ID*_fullViolin_Raw.*;

4.3.9 K-means analysis

K-means analysis is a method to examine the trend of relative quantification changes of a metabolite in different sample groups. K-means is performed based on the Z-score normalized relative quantification value.



Fig 29: K-Means diagram of differential metabolites Note: The X-axis represents the sample names and the Y-axis represents the normalized relative quantification. "Sub Class" represents a group of metabolites with the same trend and the number represent the number of metabolites in this cluster.

Figure of K-means clustering: Final report/2. Basic_analysis/kmeans/kmeans_cluster.*

4.3.10 Differential metabolite statistics

The number of different metabolites in each group is shown in the table below:

|--|

group name	All sig diff	down regulated	up regulated
A_vs_B	33	20	13
A_vs_C	31	16	15

Statistical table of differential metabolites: Final report/2.Basic_analysis/Difference_analysis/sigMetabolitesCount.xl

4.3.11 Venn diagram of differences among groups

Venn diagram was used to show the relationship between different metabolites in each group. Show petals in 5 groups or more. The results are shown below:



Fig 30: Venn diagram of differences among groups Note: Each circle in the figure represents a comparison group, the number of circles and overlapped parts represents the number of common differential metabolites between comparison groups, and the number of non-overlapped parts represents the number of unique differential metabolites in comparison groups.

/2.Basic analysis/Venn

4.4 Functional annotation and enrichment analysis of differential metabolites in KEGG database

KEGG (Kyoto Encyclopedia of Genes and Genomes) is a database that integrates compounds and genes into metabolic pathways. The KEGG database enabled researchers to study genes with their expression information and compounds with their abundances as a complete network.

4.4.1 Functional annotation of differential metabolites

Metabolites are annotated using the KEGG database, and only metabolic pathways containing differential metabolites are shown. Detailed results are found in the attached results. A portion of the results is shown below:



Fig 31: KEGG pathway of metabolites

Note: Red circles indicate that the metabolite content was significantly up-regulated in the experimental group; the blue circles indicate that the metabolite content was detected but did not change significantly; Green circles indicate that the metabolite content was significantly down-regulated in the experimental group. The orange circles indicate a mixture of both up-regulated and down-regulated metabolites. This allows searching for metabolites that may contribute to the phenotypic differences.

Final report/2.Basic_analysis/Difference_analysis/group-ID*_vs_group-ID*/enrichment/Graph/ko*.

Statistical analysis of KEGG database annotation of screened metabolites with significant differences. Some of the results are as follows:

Index	Compounds	Туре	cpd_ID
L-Aspartate	L-Aspartate	up	C00049
Glutamine	Glutamine	down	C00064
Tyrosine	Tyrosine	up	C00082
Lysine	Lysine	down	C00047
Threonine	Threonine	up	C00188
Serine	Serine	up	C00065
Itaconic-acid	Itaconic-acid	down	C00490
Malic-acid	Malic-acid	down	C03668
L-citrulline	L-citrulline	down	C00327
Dihydronicotinamide-adenine-	Dihydronicotinamide-adenine-	up	C00005
dinucleotide-phosphate(NADPH)	dinucleotide-phosphate(NADPH)		

Table	10:	KEGG	annotations	for	differential	metabolites
14010	10.	11200	annotations	101	annerentia	metholites



ko_ID	Sig_compound	compound	Sig_compound_all	compound_all
ko00220	4	9	28	59
ko00250	6	12	28	59
ko00260	4	5	28	59
ko00261	5	6	28	59
ko00270	3	5	28	59
ko00300	3	4	28	59
ko00340	2	3	28	59
ko00410	2	3	28	59
ko00470	8	12	28	59
ko00760	2	6	28	59

Table 11: Enrichment Statistics of KEGG annotations for	differential metabolites
---------------------------------------------------------	--------------------------

Final report/2.Basic_analysis/Difference_analysis/group-ID*_vs_group-ID*/enrichment/group-ID* vs_group-ID* filter kegg.xlsx.

Final report/2.Basic_analysis/Difference_analysis/group-ID*_vs_group-ID*/enrichment/group-ID* vs_group-ID* KEGG.xlsx.

4.4.2 KEGG classification of differential metabolites

The significant differential metabolites were classified based on pathway annotation . The results are as follows:



Fig 32: KEGG classification of differential metabolites Note: the Y-axis shows the name of the KEGG pathway. The number of metabolites and the proportion of the total metabolites are shown next to the bar plot.

Final report/2.Basic_analysis/Difference_analysis/group-ID*_vs_group-ID*/enrichment/group-ID*_vs_group-ID_KEGG_barplot.*.

4.4.3 Hierarchical Cluster Analysis of differential metabolites in KEGG signaling pathway

We clustered the metabolites in each pathway base on their relative quantification in order to examine the pattern of metabolite changes in different sample groups. Only pathways with at least 5 differential metabolites were analyzed.



Fig 33: Clustering heat map of differential metabolites in KEGG pathway Note: The X-axis shows the name of the samples and the Y-axis shows the differential metabolites. Different colors in the heatmap represent the values obtained after normalization and reflects the level of relative quantification. The darker the red, the higher the quantification. In contrast, the darker the green, the lower the quantification. The colored bar on top depicts sample groups. If hierarchical clustering is performed, the clustering tree will be shown on the left. If classification was performed on the metabolites, a colored bar will be shown on the left to depict classifications.

Final report/2.Basic_analysis/Difference_analysis/group-ID*_vs_group-ID*/enrichment/group-ID* vs_group-ID KEGG heatmap.*.

4.4.4 KEGG enrichment analysis of differential metabolites

KEGG pathway enrichment analysis was conducted based on the annotation results. We calculated the Rich Factor for each pathway, which is the ratio of the number of differenetial metabolites in the corresponding pathway to the total number of metabolites annotated in the same pathway. The greater the Rich Factor, the greater the degree of enrichment. P-value is the calculated using hypergeometric test as shown below:

$$P = 1 - \sum_{i=0}^{m-1} \frac{\binom{M}{i}\binom{N-M}{n-i}}{\binom{N}{n}}$$

N represents the total number metabolites with KEGG annotation, n represents the number of differential metabolites in N, M represents the number of metabolites in a KEGG pathway in N, and m represents the



number of differential metabolites in a KEGG pathway in M. The closer the p-value to 0, the more significant the enrichment. The size of the dots in the figure represents the number of significantly different metabolites enriched in the corresponding pathway. The results are shown below:



Fig 34: KEGG enrichment diagram of differential metabolites Note: The X-axis represents the Rich Factor and the Y-axis represents the pathway. The color of points reflects the p-value. The darker the red, the more significant the enrichment. The size of the dot represents the number of enriched differential metabolites.

Final report/2.Basic_analysis/Difference_analysis/group-ID*_vs_group-ID*/enrichment/group-ID* vs_group-ID* KEGG Enrichment.*.

4.4.5 Overall changes in KEGG metabolic pathway

Differential Abundance Score (DA Score) is a score based on changes in metabolites in a pathway. DA Score can capture the overall changes of all Differential metabolites in a pathway with the following formulat:

DA score=(up regulated metabolites in a pathway-down regulated metabolites in a pathway)/(Total number of metabolites annotation in a pathway)







Note: The Y-axis represents the name of differential pathway, and the X-axis represents DA Score. DA Score reflects the overall change of all metabolites in the metabolic pathway. A Score of 1 indicates that the expression trend of all identified metabolites in this pathway is up-regulated, and -1 indicates that the expression trend of all identified metabolites in this pathway is down-regulated. The length of the line represent the absolute value of DA-score while the size of the dot at the end of the line represent the number of differential metabolites. A dot on the left of the line represent the pathway is up-regulated; a dot on the right of the line represents the pathway is up-regulated. The color of the line and dot represent the p-value. The darker the red, the smaller the p-value and the darker the purple, the larger the p-value.

Final report/2.Basic Analysis/Difference analysis/group-ID* vs group-ID*/enrichment/*DA score*

4.5 Associated diseases

We annotated disease information according to the HMDB database for differential metabolites. Some of the results are shown below :



CompoundName	HmdbDiseases
L-Aspartate	Epilepsy Cirrhosis Alzheimer's disease Dicarboxylic aminoaciduria Growth hormone deficiency Schizophrenia Irritable bowel syndrome Ulcerative colitis Colorectal cancer Crohn's disease Gout Perillyl alcohol administration for cancer treatment Pancreatic cancer Periodontal
Glutamine	disease Frontotemporal dementia Lewy body disease Attachment loss Missing teeth Periodontal Probing Depth Eosinophilic esophagitis Epilepsy Schizophrenia Carbamoyl Phosphate Synthetase Deficiency Colorectal cancer Early preeclampsia Pregnancy Late-onset preeclampsia Fumarase deficiency N-acetylglutamate synthetase deficiency Obesity Glutamine deficiency, congenital Lipoyltransferase 1 Deficiency
	Phosphoenolpyruvate Carboxykinase Deficiency 1, Cytosolic Alzheimer's disease Leukemia Propionic acidemia Irritable bowel syndrome Ulcerative colitis Crohn's disease Perillyl alcohol administration for cancer treatment Pancreatic cancer Periodontal disease Frontotemporal dementia
Tyrosine	Lewy body disease Maple syrup urine disease Eosinophilic esophagitis Epilepsy Myocardial infarction Viral infection Schizophrenia Alabimar'a disease Calarattal annar Early presedemptia Presenancy
	Late-onset preeclampsia Fumarase deficiency Tyrosinemia Tyrosinemia I Hypermethioninemia Hawkinsinuria Obesity Citrullinemia type II, neonatal-onset Hypothyroidism Leukemia Irritable bowel syndrome Lucemia calificia Autiem Crahe's disease Parillel alcohel administration
	for cancer treatment Pancreatic cancer Periodontal disease Frontotemporal dementia Lewy body disease Attachment loss Missing
Lysine	teeth Periodontal Probing Depth Cachexia Eosinophilic esophagitis Refractory localization-related epilepsy Alzheimer's disease Schizophrenia Pregnancy Lysinuric protein intolerance Fumarase deficiency Histidinemia Citrullinemia type II, neonatal-onset Pyruvate
	carboxylase deficiency Lipoyltransferase 1 Deficiency Leukemia L-2-Hydroxyglutaric aciduria Crohn's disease Ulcerative colitis Irritable
	for cancer treatment Pancreatic cancer Autism Periliyi alcohol administration Frontotemporal dementia Lewy body disease Attachment loss Missing
	teeth Periodontal Probing Depth Supragingival Calculus Carbamoyl Phosphate Synthetase Deficiency Autosomal dominant polycystic kidney disease Propionic acidemia Tyrosinemia I Eosinophilic esophagitis Hyperdibasic aminoaciduria L Cystinuria Alpha-aminoacinic and
Threonine	alpha-ketoadipic aciduria 2,4-dienoyl-CoA reductase deficiency Hyperlysinuria Hyperlysinemia I, familial Epilepsy Heart failure Early preeclampsia Pregnancy Late-onset
	preeclampsia Fumarase deficiency Obesity Citrullinemia type II, neonatal-onset Pyridoxamine 5-prime-phosphate oxidase deficiency Leukemia Schizophrenia Irritable bowel syndrome Ulcerative colitis
	Colorectal cancer Crohn's disease Autism Rheumatoid arthritis Perillyl alcohol administration for cancer treatment Pancreatic cancer Periodontal disease Frontotemporal dementia Lewy body disease Attachment loss
Serine	Missing teeth Periodontal Probing Depth Alzheimer's disease Autosomal dominant polycystic kidney disease Eosinophilic esophagitis Refractory localization-related epilepsy Juvenile myoclonic epilepsy Alzheimer's disease Heart foilure Schirzophrania Lung Cancer Early
	Pregnancy Late-onset preeclampsia Sarcosinemia Fumarase deficiency Obesity Neu-Laxova Syndrome 1 Phosphoserine Aminotransferase Deficiency Phosphoserine Phosphatase Deficiency
	Leukemia 3-Phosphoglycerate dehydrogenase deficiency Serine deficiency syndrome, infantile Irritable bowel syndrome Ulcerative colitis Colorectal cancer Autism Crohn's disease Perillyl alcohol administration for cancer
	treatment Pancreatic cancer Periodontal disease Frontotemporal dementia Lewy body disease Autosomal dominant polycystic kidney disease Fosinophilic esophagitis
Itaconic-acid	
Malic-acid	Colorectal cancer Schizophrenia Anoxia Alzheimer's disease
	Frontotemporal dementia Lewy body disease Temporomandibular joint
	Cystic fibrosis Eosinophilic esophagitis 2-Ketoglutarate dehydrogenase
3	7 complex deficiency Deafness, Onychodystrophy, Osteodystrophy, Mental
L-citrulline	Retardation, and Seizures Syndrome Intestinal failure Enilensy Alzheimer's disease Argininosuccinic aciduria
	Fumarase deficiency N-acetylglutamate synthetase deficiency Citrullinemia type I Citrullinemia type II, adult-onset Pearson Syndrome

Table 12: Table of association between differential metabolites and diseases

Final report/2.Basic_analysis/Difference_analysis/group-ID*_vs_group-ID*/enrichment/group-ID*_vs_group-ID*_sigDiseasesTable.xlsx.

5 References

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6 Appendix

6.1 Analytical methods

1.PCA

Unsupervised PCA (principal component analysis) was performed by statistics function prcomp within R (www.r-project.org). The data was unit variance scaled before unsupervised PCA.

2. Hierarchical Cluster Analysis and Pearson Correlation Coefficients

The HCA (hierarchical cluster analysis) results of samples and metabolites were presented as heatmaps with dendrograms, while pearson correlation coefficients (PCC) between samples were caculated by the cor function in R and presented as only heatmaps. Both HCA and PCC were carried out by R package pheatmap. For HCA, normalized signal intensities of metabolites (unit variance scaling) are visualized as a color spectrum.

3.Differential metabolites selected

Significantly regulated metabolites between groups were determined by VIP and absolute Log_2FC (fold change). VIP values were extracted from OPLS-DA result, which also contain score plots and permutation plots, was generated using R package MetaboAnalystR. The data was log transform (log_2) and mean centering before OPLS-DA. In order to avoid overfitting, a permutation test (200 permutations) was performed.

4.KEGG annotation and enrichment analysis

Identified metabolites were annotated using KEGG compound database (http://www.kegg.jp/kegg/ compound/), annotated metabolites were then mapped to KEGG Pathway database (http://www.kegg.jp/ kegg/pathway.html). Pathways with significantly regulated metabolites mapped to were then fed into MSEA (metabolite sets enrichment analysis), their significance was determined by hypergeometric test's P-Values.

6.2 List of software and versions

Analysis	Software	Version
PCA	R (base package)	3.5.1
Pearson Correlation	R (base package; Hmisc)	3.5.1; 4.4.0
Correlation plot	R (corrplot)	0.84
Heatmap	R (heatmaply; ComplexHeatmap)	1.2.1; 2.7.1.1009
OPLS-DA	R (MetaboAnalystR)	1.0.1
Radar plot	R (fmsb)	0.7.0
Chord diagram	R (igraph; ggraph)	1.2.4.2; 2.0.2
Network diagram	R (igraph)	1.2.4.2
Regulatory network diagram	R (FELLA)	1.10.0

Table 13: Software used

Data processing methods were mainly adopted in the analysis process in two ways:

(1) unit variance scaling (UV)

Unit variance Scaling (UV) is also called Z-Score standardization, i.e., auto scaling. This method standardizes data according to mean and standard deviation of original data. The processed data conform to the standard normal distribution, that is, the mean value is 0 and the standard deviation is 1.

Calculation method: Divide the original data center by standard deviation.

The formula is as follows:

$$x'=\frac{x-\mu}{\sigma}$$

Where μ is the mean value and σ is the standard deviation.

(2) Centralization/zero-mean-centered (Ctr)

Calculation method: subtract the mean of the variables from the original data.

The formula is as follows:



$$x' = x - \mu$$