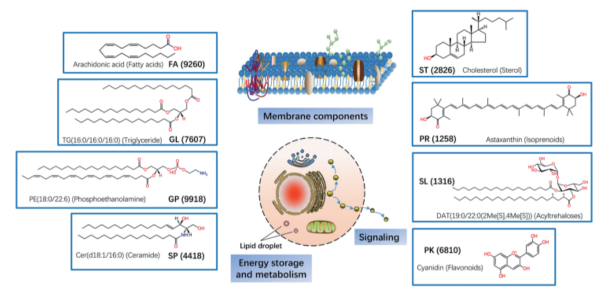


Service Description

Lipids are essential metabolites that have many key cellular functions and which can be analysed to gain insight into the metabolic state of cells. The number of lipid molecules in a cell, collectively called the lipidome, is estimated to be in the tens to hundreds of thousands.

According to the classification system proposed by the Lipid Metabolites and Pathways Strategy (LIPID MAPS) project, lipids are divided into eight classes: fatty acyls (FA), glycerolipids (GL), glycerophospholipids (GP), sphingolipids (SP), sterol lipids (ST), prenol lipids (PR), saccharolipids (SL), and polyketides (PK), covering a total of 43,413 lipid molecular species.



The common structures and main functions of eight lipid classes

Sun T, et al., Mass spectrometry-based lipidomics in food science and nutritional health: A comprehensive review, (2020)

Lipidomics is a new branch of metabolomics, which analyzes the compositions and content changes of lipids (major categories, subclasses and molecular types) in biological samples such as cells, tissues, organs or body fluids. Mass Spec based lipidomics (LC-MS), involves the comparison of the lipidome between control and test groups in order to screen differential lipids by statistical analysis, so as to identify differences between lipid metabolism and physiological/pathological changes.

BGI has extensive experience in the field of lipidomics with well-developed reliable workflows using market leading technologies and a bioinformatics infrastructure that is second to none.

Research Applications



- · Disease biomarkers research
- · Pathogenesis and prognosis study on diseases
- · Drug target research
- Animal special behavior mechanism and food/medicinal value research
- Plant growth and development research
- · Plant disease resistance and insect resistance research
- · Microbial drug resistance mechanism

Technology Platforms







Waters ACQUITY UPLC



Thermo Q Exactive/Q Exactive HF-X

Service Advantages

State-of-the-art LC-MS/MS systems

- · Thermo Q Exactive/HF
- Resolution up to 24,000, ensuring high spectral quality and accurate results

Large scale and high volume sample experience

- Sample preparation at a capacity of up to 1000+ per day
- Large scale project experience with 1000s of samples

High-precision identification results

- LipidSearch database (1.7 million lipid ions)
- 100% identification is achieved through the standards
- Identification credibility rating

Strict quality control system

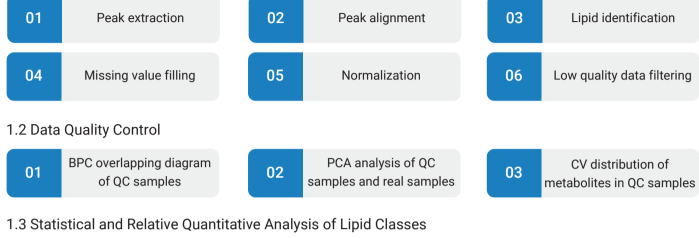
- Strict protocols governing the whole workflow
- Double quality control prcoess of isotopic internal standard and QC samples

Lipidomics Workflow

Bioinformatics Analysis Workflow

Standard:

1.1 Data Processing

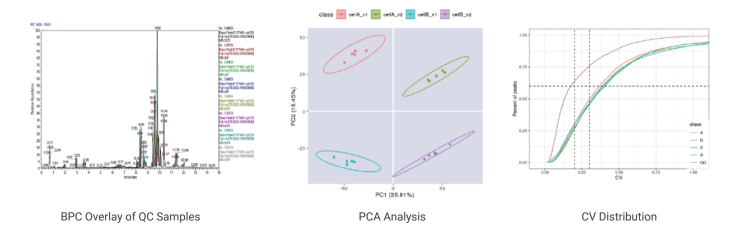


	1.4 Statistical Analysis and Screening of Differential Lipids								
	01	Univariate analysis (fold change and T test)	02	Multivariate analysis (PCA,PLS-DA)		03	Screening of differential lipids		
1.5 Differential Lipid Analysis									
	01	Volcano Plot	02	Clustering heat map		03	Lipid correlation analysis		

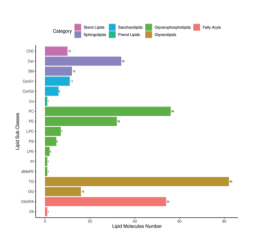
Customized Solution:

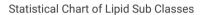
16S/Metagenome + lipidome correlation analysis Transcriptome + lipidome correlation analysis Proteome + lipidome correlation analysis

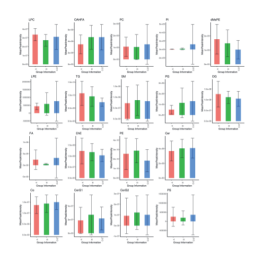
Examples of Data QC Analysis - Stability and Repeatability



Examples of Statistical and Relative Quantitative Analysis of Lipid Classes

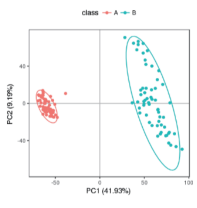




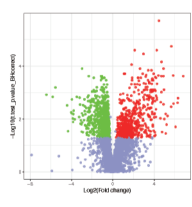


Detected Changes in Lipid Sub Class

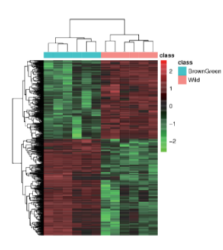
Examples of Statistical Analysis of Differential Lipids



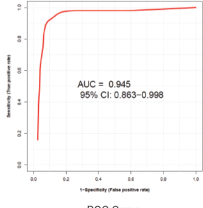
Score Graph of PLS-DA



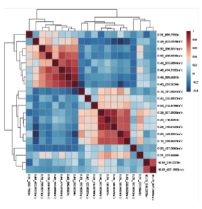
Volcano Plot



Cluster Analysis



ROC Curve



Lipid Correlation Analysis

General Sample Requirements

SAMPLE TYPE	RECOMMENDED SAMPLE AMOUNT	MINIMUM SAMPLE AMOUNT	
Serum, plasma	≥ 250 µL	≥ 50 µL	
Urine	≥ 500 µL	≥ 50 µL	
Animal and clinical tissues	≥ 200 mg	≥ 25 mg	
Feces and intestinal contents	≥ 200 mg	≥ 50 mg	
Cell	≥ 1×10 ⁷	≥ 5×10 ⁶	
Microorganism	≥ 1×10 ⁷ or ≥ 100 mg	≥ 5×10 ⁶ or ≥ 25 mg	
Culture medium, fermentation medium	≥ 1 mL	≥ 100 µL	
Plant tissue	≥ 1 g	≥ 100 mg	
Milk	≥ 1 mL	≥ 1 mL	
Other body fluids (amniotic fluid, saliva, hemolymph, cerebrospinal fluid, etc.)	≥ 250 µL	≥ 50 µL	

Turn Around Time

Sample size: 1-50, 3-5 weeks

To Learn More

To learn how your research can benefit from BGI's extensive experience in Lipidomics, visit www.bgi.com, write to us via info@bgi.com or contact your local BGI office.

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