SHUANGBIN XU

A Ph.D. student of Bioinformatics, I am broadly interested in metagenomics, bioinformatics, machine learning, data mining and visualization.

I have leaded and contributed to developing some R packages, including MicrobiotaProcess, ggtreeExtra, ggstar, ggbreak, ggtree, treeio, tidytree and ggmsa, to help related researchers to explore and understand their data.

EDUCATION current Ph.D., Bioinformaitcs

2019

2017

2016

Southern Medical University

🗣 ChuangZhou, CN

• Thesis: The development of computational methods for studying microbiome: focus on the data mining and visualization

M.S.,Aquatic Biology

JiMei University

♥ Xiamen, CN

·Thesis: 大黄鱼环状RNA挖掘与功能分析

B.S., Aquaculture
Northwest A&F University

♀ XianYang, CN

WORK EXPERIENCE

Senior Bioinformatics Engineer

Xiamen TreatGut Company

🗣 XiaMen, CN

- · Helping build the workflow of microbiome data analysis
- · Helping the mining of related microbiome data
- · Helping develop the related testing products

Bioinformatics Engineer Intern

Frasergen Genetic Information Company

• Wuhan, CN

• Helping the deployment and using with slurm and pbs scheduling systems of the software (PacBio diploid assembler: Falcon, canu, etc) of the TianHe and company's Supercomputer platform



CONTACT

- xshuangbin@163.com
- github.com/xiangpin
- BT开发者 (公众号)
- **J** (86) 13860419424

SELECTED R PACKAGES

Metagenomics

MicrobiotaProcess

Phylogeny

- · ggtreeExtra
- ggtree
- ·treeio
- tidytree

Visualization

- · ggstar
- · ggbreak

PUBLICATIONS

FIRST (AND CO-FIRST) AUTHOR

ggtreeExtra: Compact visualization of richly annotated phylogenetic data

Molecular Biology and Evolution. 2021, 38(9):4039-4042.

- · S Xu, Z Dai, P Guo, X Fu, S Liu, L Zhou, W Tang, T Feng, M Chen, L Zhan, T Wu, E Hu, Y Jiang*, X Bo*, G Yu*
- · Impact Factor₂₀₂₀ = 16.24
- Use ggbreak to effectively utilize plotting space to deal with large datasets and outliers

Frontiers in Genetics. 2021, 12:774846.

- · S Xu#, M Chen#, T Feng, L Zhan, L Zhou, G Yu*
- · Impact Factor₂₀₂₀ = 4.599

Gut microbiota and metabolite alterations associated with reduced bone 2020 mineral density or bone metabolic indexes in postmenopausal osteoporosis

Aging (Albany NY). 2020, 12(9):8583-8604.

- · J He#, **S** Xu#, B Zhang#, C Xiao, Z Chen, F Si, J Fu, X Lin, G Zheng*, G Yu*, J Chen*
- \cdot Impact Factor₂₀₂₀ = 5.682
- Leveraging Fecal Bacterial Survey Data to Predict Colorectal Tumors Frontiers in Genetics. 2019, 10:447.
 - · B Zhang", S Xu", W Xu", Q Chen, Z Chen, C Yan, Y Fan, H Zhang, Q Liu, J Yang, J Yang, C Xiao, H Xu*, J Ren*
 - \cdot Impact Factor₂₀₁₉ = 3.258
 - Transcriptome-wide identification and functional investigation of circular RNA in the teleost large yellow croaker (Larimichthys crocea)

Marine genomics. 2017, 32:71-78

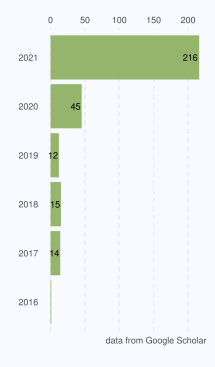
- · S Xu, S Xiao, C Qiu, Z Wang*
- Impact Factor₂₀₁₇ = 1.937
- CONTRIBUTING AUTHOR

clusterProfiler 4.0: A universal enrichment tool for interpreting omics data

The Innovation. 2021, 2(3):100141.

• T Wu#, E Hu#, S Xu, M Chen, P Guo, Z Dai, T Feng, L Zhou, W Tang, L Zhan, X Fu, S Liu, X Bo*, G Yu*

- \cdot Citation = 310
- · H-index = 8
- \cdot I10-index = 7



2021

2019

2017

2021

2020

treeio: an R package for phylogenetic tree input and output with richly annotated and associated data

Molecular Biology and Evolution. 2020, 37(2):599-603.

- · LG Wang, TTY Lam, **S Xu**, Z Dai, L Zhou, T Feng, P Guo, CW Dunn, BR Jones, T Bradley, H Zhu, Y Guan, Y Jiang, G Yu*
- · Impact Factor₂₀₂₀ = 16.24
- Featured Gut Microbiomes Associated With the Progression of Chronic Hepatitis B Disease

Frontiers in microbiology. 2020, 11:383.

- · Z Chen#, Y Xie#, F Zhou, B Zhang, J Wu, L Yang, **S Xu**, R Stedtfeld, Q Chen, J Liu, X Zhang*, H Xu*, J Ren*
- · Impact Factor₂₀₂₀ = 5.64
- Preservation of the fecal samples at ambient temperature for microbiota analysis with a cost-effective and reliable stabilizer EffcGut

Science of The Total Environment. 2020, 741:140423.

- · L Yang*, K Hou*, B Zhang*, C Ouyang, A Lin, **S Xu**, D Ke, L Fang, Q Chen, J Wu, C Yan, Y Lian, T Jiang, J He, H Wang, Y Fu, C Xiao, Z Chen*
- Impact Factor₂₀₂₀ = 7.963

2018

Genome-wide association study using single marker analysis and Bayesian methods for the gonadosomatic index in the large yellow croaker

Aquaculture. 2018, 486:26-30.

- · Y Gao[#], L Dong[#], **S Xu**, S Xiao, M Fang, Z Wang^{*}
- \cdot Impact Factor₂₀₁₈ = 3.022

2017

 Cloning and expression profiling of gsdf and amh genes in large yellow croaker (Larimichthys crocea)

South China Fisheries Science. 2017, 13(6):1-13.

- · A Lin, Y Xie, S Xu, K Ye, S Gong, Z Wang*
- Identification of a male-specific DNA marker in the large yellow croaker (Larimichthys crocea)

Aquaculture. 2017, 480:116-122.

- \cdot A Lin, S Xiao, **S Xu**, K Ye, X Lin, S Sun, Z Wang*
- \cdot Impact Factor₂₀₁₇ = 2.71

2015

Rapid construction of genome map for large yellow croaker (Larimichthys crocea) by the whole-genome mapping in BioNano Genomics Irys system

BMC Genomics. 2015, 16(1):1-10

- · S Xiao, J Li, F Ma, L Fang, **S Xu**, W Chen, Y Wang*
- \cdot Impact Factor₂₀₁₅ = 3.867



Analysis and visualization of microbiome datasets using MicrobiotaProcess

The 10th BiocAsia2020 Conference, 2020

♥ Virtual Conference

2020