Guangchuang Yu

Professor

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EDUCATION

The University of Hong Kong	Ph.D. Evolution Study of Influenza A virus	2017
Anhui Medical University	M.S. Biochemistry and Molecular Biology	2009
South China Agricultural University	B.S. Biotechnology	2005
延安干部培训学院	南方医科大学高层次人才"不忘初心 牢记使命"学习教育	2018
Institut Pasteur, Paris	Train the Trainers	2015

■ CERTIFICATE

Certificate of Teaching and Learning in Higher Education, Centre for Enhancement of Teaching and Learning, The University of Hong Kong

m EMPLOYMENT

Institute of Bioinformatics, School of Basic Medical Sciences, Southern Medical University:

Professor 2018 -

State Key Laboratory of Infectious Diseases, School of Public Health, The University of Hong Kong:

Post-doctoral Fellow 2017-2018

Key Laboratory of Functional Protein Research of Guangdong Higher Education Institutes, College of Life Science and Technology, Jinan University:

2009 - 2013Bioinformatician

SCHOLARSHIPS & AWARDS

• 南方医科大学高层次人才引进(第三层次)

- 2018
- Travel Grant for the 3rd South Lake Innovation Forum for International Young Talents, awarded by Huazhong Agricultural University
- Travel Grant for Phylogenetic tree visualization workshop/hackathon, awarded by The Field Museum of Natural History, USA
- Travel Grant for the 2nd International Workshop on Environmental Microbiomes, awarded by Sun Yat-sen University
- Postgraduate Studentships, awarded by The University of Hong Kong 2013 - 2017
- Travel Grant for Options IX for the control of influenza, awarded by Areas of Excellence Scheme (AoE) and Theme-based Research Scheme (TRS), Hong Kong 2016
- $\bullet\,$ Travel Grant for the $9^{\rm th}$ China-R conference, awarded by Capital of Statistics 2016
- Best paper of the year 2010, awarded by Journal of Integrated OMICS

2012

■ GRANTS

- (在研) PI, 南方医科大学第三层次人才引进科研项目,南方医科大学 (c1051022), CNY ¥1,500,000 2018 - 2019
- (完成) \mathbf{PI} , hnRNPK 蛋白在调控人肺癌细胞周期胞质分裂中的作用研究, 中央高校基本科研业务费专项资金 (21613414), CNY ¥100.000 2013
- (完成) PI, Hela 细胞周期动态蛋白质组学分析,中央高校基本科研业务费专项资金(21611303), $CNY\ Y20.000$

₩ TEACHING

Southern Medical University

基因组学 2019 2019

基础医学新进展讲座

University of Southern Denmark

BMB209: Workshops in Applied Bioinformatics 2019

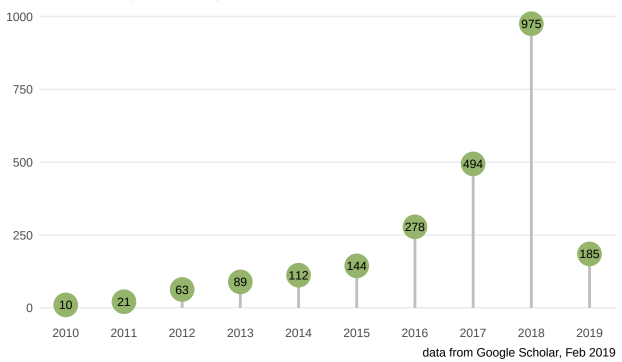
Jinan University

功能蛋白质研究 2011 - 2012

PUBLICATIONS

Journal Articles

Citation = 2410, H-index = 15, I10-index = 20



- **★** ESI Hot Paper (X1)
- TESI Highly Cited Paper (X4)
- # indicates equal contribution, * indicates corresponding
- 1. G Yu. clusterProfiler: universal enrichment tool for functional and comparative study. bioRxiv. 2018, https://doi.org/10.1101/256784
- 2. **G** Yu*, TTY Lam, H Zhu, Y Guan*. Two methods for mapping and visualizing associated data on phylogeny using ggtree. Molecular Biology and Evolution. 2018, 35(12):3041-3043. Impact Factor: 10.217
- 3. G Yu. Using meshes for MeSH term enrichment and semantic analyses. Bioinformatics. 2018, 34(21):3766-3767. Impact Factor: 5.481
- 4. \mathbf{G} \mathbf{Yu} , DK Smith, H Zhu, Y Guan, TTY Lam*. ggtree: an R package for visualization and annotation of phylogenetic trees with their covariates and other associated data. Methods in Ecology and

- 5. **G Yu**, QY He*. ReactomePA: an R/Bioconductor package for reactome pathway analysis and visualization. *Molecular BioSystems*. 2016, 12(2):477-479. ♥ Citation: 127, Impact Factor: 2.759
- 6. **G** Yu*, LG Wang, QY He*. ChIPseeker: an R/Bioconductor package for ChIP peak annotation, comparision and visualization. *Bioinformatics*. 2015, 31(14):2382-2383. ♥ Citation: 206, Impact Factor: 5.481
- 7. **G** Yu*, LG Wang, GR Yan, QY He*. DOSE: an R/Bioconductor package for Disease Ontology Semantic and Enrichment analysis. *Bioinformatics*. 2015, 31(4):608-609. Citation: 110, Impact Factor: 5.481
- 8. X Sun[#], **G** Yu[#], Q Xu, N Li, C Xiao, X Yin, K Cao, J Han, QY He^{*}. Putative cobalt- and nickel-binding proteins and motifs in *Streptococcus pneumonia*. *Metallomics*. 2013, 5(7):928-935. Citation: 19, Impact Factor: 4.069
- 9. YY Zhu[#], **G Yu**[#], Y Zhang, Z Xu, YQ Huang, GR Yan^{*}, QY He^{*}. A novel andrographolide derivative AL-1 exerts its cytotoxicity on K562 cells through a ROS-dependent mechanism. *Proteomics*, 2013, 13(1):169-178. Citation: 21, Impact Factor: 3.532
- 10. **G** Yu[#], LG Wang[#], XH Meng, QY He^{*}. LXtoo: an integrated live Linux distribution for the bioinformatics community. *BMC Research Notes*. 2012, 5(1):360. Citation: 6
- 11. **G Yu**, LG Wang, Y Han, QY He*. clusterProfiler: an R package for comparing biological themes among gene clusters. *OMICS: A Journal of Integrative Biology*. 2012, 16(5):284-287. Citation: 928, Impact Factor: 2.370
- 12. **G Yu**, QY He*. Functional similarity analysis of human virus-encoded miRNAs. *Journal of Clinical Bioinformatics*. 2011, 1(1):15. Citation: 16
- 13. **G** Yu[#], CL Xiao[#], X Bo, CH Lu, Y Qin, S Zhan, QY He^{*}. A new method for measuring functional similarity of microRNAs. *Journal of Integrated OMICS*. 2011, 1(1):49-54. Citation: 21
- 14. **G** Yu[#], CL Xiao[#], CH Lu[#], HT Jia, F Ge, W Wang, XF Yin, HL Jia, JX He^{*}, QY He^{*}. Phosphoproteome profile of human lung cancer cell line A549. *Molecular BioSystems*. 2011, 7(2):472-479. Citation: 13, Impact Factor: 2.759
- 15. **G Yu**[#], F Li[#], Y Qin, X Bo*, Y Wu, S Wang*. GOSemSim: an R package for measuring semantic similarity among GO terms and gene products. *Bioinformatics*. 2010, 26(7):976-978. Citation: 423, Impact Factor: 5.481
- 16. XY Yang, K He, G Du, X Wu, G Yu, Y Pan, G Zhang*, X Sun*, QY He*. Integrated Translatomics with Proteomics to Identify Novel Iron-Transporting Proteins in *Streptococcus pneumoniae*. *Frontiers in Microbiology*. 2016, 7:78. Citation: 15, Impact Factor: 4.019
- 17. PY Huang, CCD Lee, CH Yip, CL Cheung, **G Yu**, TTY Lam, DK Smith, H Zhu*, Y Guan*. Genetic characterization of highly pathogenic H5 influenza viruses from poultry in Taiwan, 2015. *Infection, Genetics and Evolution*. 2016, 38:96-100. Citation: 8, Impact Factor: 2.545
- 18. SH Xu, JZ Huang, ML Xu, G Yu, XF Yin, D Chen*, GR Yan*. ACK1 Promotes Gastric Cancer Epithelial-Mesenchymal Transition and Metastasis through AKT-POU2F1-ECD Signaling. *Journal of Pathology*. 2015, 236(2):175-185. Citation: 27, Impact Factor: 6.253
- 19. Z Guo, J Han, XY Yang, K Cao, K He, G Du, G Zeng, L Zhang, **G Yu**, Z Sun, QY He^{*}, X Sun^{*}. Proteomic analysis on the copper resistance of *Streptococcus pneumonia*. *Metallomics*. 2015, 7(3):448-454. Citation: 11, Impact Factor: 4.069
- 20. XY Yang, L Zhang, J Liu, N Li, **G Yu**, K Cao, J Han, G Zeng, Y Pan, X Sun*, QY He*. Proteomic analysis on the antibacterial activity of a Ru(II) complex against *Strepococcus pneumonia*. *Journal of Proteomics*. 2015, 115:107-116. Citation: 11, Impact Factor: 3.722
- 21. N Wang[#], J Zhang[#], L Zhang, XY Yang, N Li, **G Yu**, J Han, K Cao, Z Guo, X Sun^{*}, QY He^{*}. Proteomic analysis of putative heme-binding proteins in *Streptococcus pyogenes*. *Metallomics*. 2014, 6(8):1451-1459. Citation: 0, Impact Factor: 4.069
- 22. H Sun*, J Weng, G Yu, RH Massawe. A DNA-Based Semantic Fusion Model for Remote Sensing Data. *PLoS One.* 2013, 8(10):e77090. Citation: 6, Impact Factor: 2.766
- 23. GR Yan[#], Z Tan[#], Y Wang, ML Xu, **G Yu**, Y Li^{*}, QY He^{*}. Quantitative Proteomics Characterization on the Anti-tumor Effects of Isodexyelephantopin against Nasopharyngeal Carcinoma. *Proteomics*. 2013, 13(21):3222-3232. Citation: 16, Impact Factor: 3.532
- 24. J Li, X Lu, Q Wu, G Yu, Z Xu, L Qiu, Z Pei, Y Lin, J Pang*. Design, SAR, angiogenic activities evaluation and pro-angiogenic mechanism of new marine cyclopeptide analogs. *Current Medicinal*

- Chemistry. 2013, 20(9):1183-1194. Citation: 3, Impact Factor: 3.469
- 25. Y Han, **G Yu**, H Sarioglu, A Caballero-Martinez, F Schlott, M Ueffing, H Haase, C Peschel, AM Krackhardt*. Proteomic investigation of the interactome of FMNL1 in hematopoietic cells unveils a role in calcium-dependent membrane plasticity. *Journal of Proteomics*. 2013, 78:72-82. Citation: 9, Impact Factor: 3.722
- 26. Y Ma[#], J Yang[#], X Fan[#], H Zhao, W Hu, Z Li, **G Yu**, X Ding, J Wang, X Bo, X Zheng, Z Zhou^{*}, S Wang^{*}. Cellular MicroRNA let-7c Inhibits M1 Protein Expression of the H1N1 Influenza A Virus in Infected Human Lung Epithelial Cells. *Journal of Cellular and Molecular Medicine*. 2012, 16(10):2539-2546. Citation: 61, Impact Factor: 4.302
- 27. GR Yan[#], FY Zhou[#], BL Dang, Y Zhang, **G Yu**, X Liu, QY He^{*}. Genistein-induced mitotic arrest of gastric cancer cells by downregulating KIF20A, a proteomics study. *Proteomics*. 2012, 12(14):2391-2399. Citation: 52, Impact Factor: 3.532
- 28. X Sun[#], XY Yang[#], XF Yin, **G Yu**, CL Xiao, X He, QY He^{*}. Proteomic Analysis of Membrane Proteins from *Streptococcus pneumoniae* with Multiple Separation Methods plus High Accuracy Mass Spectrometry. *OMICS: A Journal of Integrative Biology*. 2011, 15(10):683-694. Citation: 16, Impact Factor: 2.370
- 29. **余光创**,王超,许尹,伯晓晨,王升启^{*},秦宜德^{*}. 结合基因组分析预测炎症相关 miRNA 及其靶标. **军事医学科学院院刊**. $2009,\ 33(3):245-247.$
- 30. 余光创,秦宜德,伯晓晨,王升启 * . 依赖于 5^* 端非编码区高级结构的真核生物 mRNA 翻译调控. 中国生物化学与分子生物学报. $2007,\ 23(11):881-887.$

Books

• **余光创**, "生物信息学 II", 《功能蛋白质研究》, 385-425, 何庆瑜 (主编), **科学出版社**, 2012. ("十二五"国家重点图书出版 规划项目)

SOFTWARE DEVELOPMENT

R ranking: ranked 30/72305 worldwide by Git Awards

- badger: Badge for R Package
- dlstats: Download Stats of R Packages
- emojifont: Emoji and Font Awesome in Graphics
- ggplotify: Convert Plot to 'grob' or 'ggplot' Object
- ggimage: Use Image in 'ggplot2'
- meme: Create Meme
- rvcheck: R/Package Version Check
- scatterpie: Scatter Pie Plot
- shadowtext: Shadow Text Grob and Layer
- ChIPseeker: ChIPseeker for ChIP peak Annotation, Comparison, and Visualization
- clusterProfiler: statistical analysis and visualization of functional profiles for genes and gene clusters
- DOSE: Disease Ontology Semantic and Enrichment analysis
- enrichplot: Visualization of Functional Enrichment Result
- ggtree: an R package for visualization and annotation of phylogenetic trees with their covariates and other associated data
- GOSemSim: GO-terms Semantic Similarity Measures
- meshes: MeSH Enrichment and Semantic analyses
- Reactome Pathway Analysis
- seqcombo: Visualization Tool for Sequence Recombination and Reassortment
- tidytree: A Tidy Tool for Phylogenetic Tree Data Manipulation
- TIPars: Taxa Insertion by Parsimony
- treeio: Base Classes and Functions for Phylogenetic Tree Input and Output

Contributions to open source libraries

- basicTrendline: Add Trendline and Confidence Interval of Basic Regression Models to Plot
- cowsay: Messages, Warnings, Strings with Ascii Animals

- kableExtra: Construct Complex Table with 'kable' and Pipe Syntax
- microbiomeViz: Visualizing Microbiome Data
- MSnbase: Base Functions and Classes for Mass Spectrometry and Proteomics
- pheatmap: Pretty Heatmaps
- RIdeogram: Visualize and Map Genome Wide Data in Ideograms
- scholar: Analyse Citation Data from Google Scholar

♠ PROFESSIONAL AFFILIATIONS

• Member, International Society for Influenza and other Respiratory Virus Disease

2016 2015

• Steering Committee, International Network for Data Analysis, Institut Pasteur

• Visiting student, Beijing Institute of Radiation Medicine

2005-2009

▲ PROFESSIONAL SERVICES

Associate Editor

• Journal of Integrated OMICS, 2011-present

Reviewer

Bioinformatics, BMC Bioinformatics, Computational Biology and Chemistry, Current Bioinformatics, Gastroenterology Research and Practice, Molecular BioSystems, Nucleic Acids Research, R Journal

■ PUBLIC MEDIA

• "Plotting trees + data" lecture by Guangchuang Yu. https://youtu.be/tqspAXM0UQE

CONFERENCE PROCEEDINGS

Presentations

- 1. Two methods for mapping and visualizing associated data on phylogeny using ggtree, 第一届南方医科大学 基础医学学术年会, 南方医科大学, 广州, 2019 年 1 月
- 2. Data integration and visualization of phylogenetic trees, The 3rd South Lake Innovation Forum for International Young Talents, Huazhong Agricultural University, Wuhan, May, 2018
- 3. getree for tree visualization & annotation (invited), Phylogenetic tree visualization workshop/hackathon, The Field Museum of Natural History, Chicago, May. 2017
- 4. Computational Methods in Ecology and Evolution, The Second International Workshop on Environmental Microbiomes, Sun Yat-sen University, Guangzhou, April. 2017
- 5. ggtree for visualization and annotation of phylogenetic trees (invited), The 9th China-R Conference, Renmin University of China, Beijing, May. 2016

Posters

- 1. ggtree: a programming package for complex data integration and analysis in phylogeny to identify potential genetic markers of influenza a virus that determine interspecies transmission, Scientific Advisory Board (SAB) Meeting for the Theme-based Research Scheme on Viral, Host and Environmental Determinants of Influenza Virus Transmission and Pathogenesis, The University of Hong Kong, Hong Kong, Jun. 2017
- 2. Phylogeny-quided genome assembly method for short read nucleotide sequences from co-infecting influenza A viruses, Options IX for the control of influenza, International Society for Influenza and other Respiratory Virus Disease, Chicago, Aug. 2016
- 3. ggtree: an R package for visualization and annotation of phylogenetic tree with different types of meta-data. The 20th Research Postgraduate Symposium. The University of Hong Kong, Hong Kong, Dec. 2015