# Guangchuang Yu

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**2013-2017 PhD**; The University of Hong Kong; *Evolution Study of Influenza A Virus* 

**2006-2009 MSc**; Anhui Medical University; *Biochemistry and Molecular Biology* 

**2001-2005 BSc**; South China Agricultural University; *Biotechnology* 

**2015** Institut Pasteur, Paris; *Train the Trainers 2015* 

### **Professional Experience**

**2016 Member**; International Society for Influenza and other Respiratory Virus Diseases

2015 INDA Steering Committee; International Network for Data Analysis, Institut Pas-

teur

**2009-2013** Bioinformatician; College of Life Science and Technology, Jinan University

**2007-2009 Visiting Student**; Beijing Institute of Radiation Medicine

# **Teaching Experience**

2013-2014 Student Helper; BIOL6001 Presentation Skills and Research Seminars in Biolog-

ical Sciences (postgraduate course), The University of Hong Kong

**2011-2012** Lecturer; Functional Proteomics (postgraduate course), Jinan University

#### Awards

**2013-2017** Postgraduate Studentships (for PhD students at The University of Hong Kong)

2012 Best paper of the year 2010, Journal of Integrated OMICS.

#### Grants

2013 (completed) PI, Investigate functions of hnRNPK in regulating cytokinesis of hu-

man lung cancer cell, Fundamental Research Funds for the Central Universities

(21613414), 100,000

(completed) PI, Dynamic proteomic analysis of HeLa cell cycle, Fundamental Re-

search Funds for the Central Universities (21611303), 20,000

### **Professional Services**

**Associate** 

Journal of Integrated OMICS, 2011-

**Editor** 

Reviewer Bi

**Bioinformatics** 

**BMC Bioinformatics** 

Computational Biology and Chemistry

**Current Bioinformatics** 

Gastroenterology Research and Practice

Molecular BioSystems

R Journal

# Languages

Teochew Native language

Chinese Man-

Very fluent

darin

Cantonese

Very fluent

**English** fluent

# Programming Skills

#### R

- More than 8 years of experience
- Developed 14 packages (6 on CRAN and 8 on Bioconductor)
- Ranked 64/60358 worldwide by Git Awards

#### Unix

- More than 14 years of experience (Linux & BSD)
- Experience of server setup, management and application deployment

#### Other

- Good knowledge of Git, LATEX Bash, C/C++ and Java
- Basic knowledge of Perl, Python and Matlab

# Software Development

R

badger Badge for R Package

Monthly download stats of 'CRAN' and 'Bioconductor' packages distats

emojifont Using emoji font in both 'base' and 'ggplot2' graphics

Use images in 'ggplot2' graphic system ggimage

rvcheck Check latest release version of R and R package

scatterpie Creates scatterpie plots, especially useful for plotting pies on a map

#### **Bioconductor**

**ChIPseeker** ChIP peak Annotation, Comparison and Visualization

clusterPro-Statistical analysis and visualization of functional profiles for genes and gene clus-

filer ters

**DOSE** Disease Ontology Semantic and Enrichment analysis

Visualization and annotation of phylogenetic trees with associated data ggtree

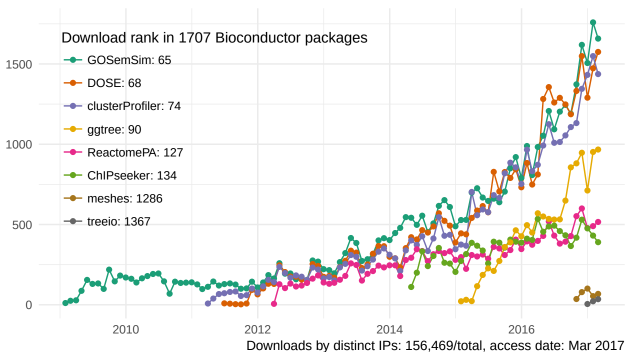
**GOSemSim** GO and gene samantic similarity measurement

meshes MeSH enrichment and semantic analyses

ReactomePA Reactome pathway analysis

treeio Base Classes and Functions for Phylogenetic Tree Input and Output

### Monthly download stats



#### Java

**TIPars** Taxa Insertion by Parsimony

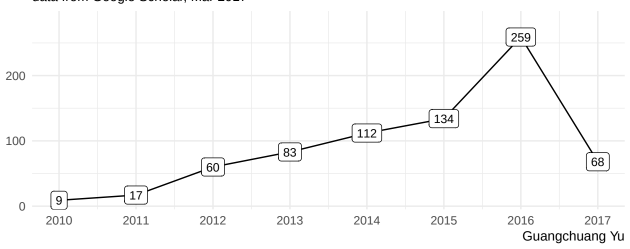
# Contributions to open source libraries

cowsay Messages, Warnings, Strings with Ascii Animals

MSnbase Base Functions and Classes for MS-based Proteomics

# Research Highlights

Citation = 772, H-index = 12, I10-index = 16 data from Google Scholar, Mar 2017



- 25 peer-reviewed publications (IF5 = 5-year Impact Factor)
  - Methods in Ecology of Evolution (X1, rank 11 in Environment Ecology, IF5 = 8.121)
  - Bioinformatics (X3, rank 4 in Computer Science, IF5 = 7.685)
  - Frontiers in Microbiology (X1, rank 21 in Microbiology, IF5 = 4.36)
  - Proteomics (X3)
  - Journal of Proteomics (X2)
  - Metallomics (X3)
  - Molecular BioSystems (X2)
- 12 peer-reviewed publications of first author
- 4 papers of first author with IF5 >= 5
- 2 papers of first author with citations >= 100
- view my publication impact on Impactstory

#### **Ontology Semantic Similarity Analysis**

- developed 3 packages: GOSemSim, DOSE and meshes
- propose a method for measuring functional similarity of microRNAs and applied it to analyze human virus-encoded miRNAs
- published 4 papers, including 2 in *Bioinformatics*

#### **Pathway Analysis**

- developed 2 packages: clusterProfiler and ReactomePA
- published 2 papers in OMICS: A Journal of Integrative Biology and Molecular BioSystems

#### **Genomic Analysis**

- developed ChIPseeker for ChIP-seq data annotation, visualization and data mining (by integrated GEO database)
- published 1 paper in Bioinformatics

#### **Proteomic Analysis**

- contributed to open source library, *MSnbase*, developed by Laurent Gatto (University of Cambridge)
- published 9 papers in Molecular BioSystems, Metallomics, Proteomics and Journal of Proteomics

#### **Evolutionary Analysis**

- developed ggtree for parsing tree files and supporting tree visualization and annotation by grammar of graphics
- devleloping phylogeny-guided genome assembly method for short read nucleotide sequences from co-infecting influenza viruses
- published 1 paper in Methods in Ecology and Evolution

### Publications

- 1. **G 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 Evolution***. 2017, 8(1):28-36.**
- 2. **G Yu**, QY He<sup>\*</sup>. ReactomePA: an R/Bioconductor package for reactome pathway analysis and visualization. *Molecular BioSystems*. 2016, 12(2):477-479.
- 3. **G Yu**\*, LG Wang, QY He\*. **ChIPseeker: an R/Bioconductor package for ChIP peak annotation, comparision and visualization. Bioinformatics**. 2015, 31(14):2382-2383.
- 4. **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.
- 5. **G** Yu<sup>†</sup>, LG Wang<sup>†</sup>, XH Meng, QY He<sup>\*</sup>. **LXtoo: an integrated live Linux distribution for the bioinformatics community.** *BMC Research Notes***. 2012, 5(1):360.**
- G Yu, LG Wang, Y Han, QY He<sup>\*</sup>. clusterProfiler: an R package for comparing biological themes among gene clusters. *OMICS: A Journal of Integrative Biology*. 2012, 16(5):284-287.
- 7. **G Yu**, QY He<sup>\*</sup>. Functional similarity analysis of human virus-encoded miRNAs. *Journal of Clinical Bioinformatics*. 2011, 1(1):15.
- 8. **G Yu**<sup>†</sup>, CL Xiao<sup>†</sup>, X Bo, CH Lu, Y Qin, S Zhan, QY He<sup>\*</sup>. **A new method for measuring functional similarity of microRNAs**. *Journal of Integrated OMICS*. 2011, 1(1):49-54.
- G Yu<sup>†</sup>, CL Xiao<sup>†</sup>, CH Lu<sup>†</sup>, HT Jia, F Ge, W Wang, XF Yin, HL Jia, JX He<sup>\*</sup>, QY He<sup>\*</sup>. Phosphoproteome profile of human lung cancer cell line A549. Molecular BioSystems. 2011, 7(2):472-479.
- 10. **G** Yu<sup>†</sup>, F Li<sup>†</sup>, Y Qin, X Bo<sup>\*</sup>, Y Wu, S Wang<sup>\*</sup>. **GOSemSim: an R package for measuring semantic similarity among GO terms and gene products.** *Bioinformatics***. 2010, 26(7):976-978.**
- 11. X Sun<sup>†</sup>, **G Yu**<sup>†</sup>, Q Xu, N Li, C Xiao, X Yin, K Cao, J Han, QY He<sup>\*</sup>. **Putative cobalt- and nickel-binding proteins and motifs in** *Streptococcus pneumonia*. *Metallomics*. 2013, 5(7):928-935.

- 12. YY Zhu<sup>†</sup>, **G Yu**<sup>†</sup>, Y Zhang, Z Xu, YQ Huang, GR Yan<sup>\*</sup>, QY He<sup>\*</sup>. **A novel andrographolide derivative AL-1 exerts its cytotoxicity on K562 cells through a ROS-dependent mechanism.** *Proteomics*, 2013, 13(1):169-178.
- 13. XY Yang, K He, G Du, X Wu, **G Yu**, Y Pan, G Zhang<sup>\*</sup>, X Sun<sup>\*</sup>, QY He<sup>\*</sup>. **Integrated Translatomics with Proteomics to Identify Novel Iron–Transporting Proteins in** *Streptococcus pneumoniae*. *Frontiers in Microbiology*. 2016, 7:78.
- 14. PY Huang, CCD Lee, CH Yip, CL Cheung, **G Yu**, TTY Lam, DK Smith, H Zhu<sup>\*</sup>, Y Guan<sup>\*</sup>. **Genetic** characterization of highly pathogenic H5 influenza viruses from poultry in Taiwan, 2015. *Infection, Genetics and Evolution*. 2016, 38:96-100.
- 15. SH Xu, JZ Huang, ML Xu, **G Yu**, XF Yin, D Chen<sup>\*</sup>, GR Yan<sup>\*</sup>. **ACK1 Promotes Gastric Cancer Epithelial-Mesenchymal Transition and Metastasis through AKT-POU2F1-ECD Signaling.** *The Journal of Pathology*. 2015, 236(2):175-185.
- 16. Z Guo, J Han, XY Yang, K Cao, K He, G Du, G Zeng, L Zhang, **G Yu**, Z Sun, QY He<sup>\*</sup>, X Sun<sup>\*</sup>. **Proteomic analysis on the copper resistance of** *Streptococcus pneumonia. Metallomics*. 2015, 7(3):448-454.
- 17. XY Yang, L Zhang, J Liu, N Li, **G Yu**, K Cao, J Han, G Zeng, Y Pan, X Sun<sup>\*</sup>, QY He<sup>\*</sup>. **Proteomic** analysis on the antibacterial activity of a Ru(II) complex against *Strepococcus pneumonia*. *Journal of Proteomics*. 2015, 115:107-116.
- 18. N Wang<sup>†</sup>, J Zhang<sup>†</sup>, L Zhang, XY Yang, N Li, **G Yu**, J Han, K Cao, Z Guo, X Sun<sup>\*</sup>, QY He<sup>\*</sup>. **Proteomic analysis of putative heme-binding proteins in** *Streptococcus pyogenes*. *Metallomics*. 2014, 6(8):1451-1459.
- 19. H Sun, J Weng, G Yu, RH Massawe. A DNA-Based Semantic Fusion Model for Remote Sensing Data. *PLoS One*. 2013, 8(10):e77090.
- 20. GR Yan<sup>†</sup>, Z Tan<sup>†</sup>, Y Wang, ML Xu, **G Yu**, Y Li<sup>\*</sup>, QY He<sup>\*</sup>. **Quantitative Proteomics Characterization on the Anti-tumor Effects of Isodexyelephantopin against Nasopharyngeal Carcinoma.** *Proteomics***. 2013, 13(21):3222-3232.**
- 21. J Li, X Lu, Q Wu, **G Yu**, Z Xu, L Qiu, Z Pei, Y Lin, J Pang<sup>\*</sup>. **Design, SAR, angiogenic activities** evaluation and pro-angiogenic mechanism of new marine cyclopeptide analogs. *Current Medicinal Chemistry*. 2013, 20(9):1183-1194.
- 22. Y Han, **G Yu**, H Sarioglu, A Caballero-Martinez, F Schlott, M Ueffing, H Haase, C Peschel, AM Krackhardt<sup>\*</sup>. **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.
- 23. Y Ma<sup>†</sup>, J Yang<sup>†</sup>, X Fan<sup>†</sup>, H Zhao, W Hu, Z Li, **G Yu**, X Ding, J Wang, X Bo, X Zheng, Z Zhou<sup>\*</sup>, S Wang<sup>\*</sup>. 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.
- 24. GR Yan<sup>†</sup>, FY Zhou<sup>†</sup>, BL Dang, Y Zhang, **G Yu**, X Liu, QY He<sup>\*</sup>. **Genistein-induced mitotic arrest of gastric cancer cells by downregulating KIF20A**, a proteomics study. *Proteomics*. 2012, 12(14):2391-2399.
- 25. X Sun<sup>†</sup>, XY Yang<sup>†</sup>, XF Yin, **G Yu**, CL Xiao, X He, QY He<sup>\*</sup>. 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.

### Book Chapters

1. **G Yu**, "Bioinformatics II", in "Functional Protein Research", 385-425, Qing-Yu He (*Eds*), *Science Press*, 2012. (China's Twelfth Five-Year National Important Books Project, in Chinese)

### Conference Presentations

1. **ggtree for visualization and annotation of phylogenetic trees** (invited), The 9<sup>th</sup> China-R Conference, Renmin University of China, Beijing, May. 2016

### Conference Posters

- 1. Phylogeny-guided 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
- 2. **ggtree:** an *R* package for visualization and annotation of phylogenetic tree with different types of meta-data, The 20<sup>th</sup> Research Postgraduate Symposium, The University of Hong Kong, Hong Kong, Dec. 2015