# RUI FU

#### RNA Bioscience Initiative, University of Colorado Anschutz Medical Campus

My research interests are in RNA-binding protein-mediated post-transcriptional gene expression regulation, particularly RNA decay, interrogated through biochemical experiments and computational analyses of genomic data.



## **EDUCATION**

2015 2009

2009 B.S. Biological Science Xiamen University 2005

Ph.D. Biological Science University of California San Diego

Q La Jolla, CA, USA

• Xiamen, Fujian, China



### RESEARCH EXPERIENCE

Current 2017

## Semi-Independent Postdoctoral Informatics Fellow

Jay Hesselberth's Supervision

- **Q** University of Colorado Anschutz
- · Development of bioinformatics software for single cell RNA sequencing
- · RNA dynamics modeling of steroidogenesis
- · RNA-seg analysis and transcriptome annotation improvements for hibernating ground squirrel brain and liver
- · Collaboration on various single cell RNA sequencing projects with clinical investigators
- · Analysis of human genetic variation from gnomAD database for insights into nonsense-mediated mRNA decay

2017 2016

#### Postdoctoral Research Fellow

Judy Lieberman's Lab

• Harvard Medical School

- · Identification of a mitochondrial trigger for apoptotic mRNA decay
- · Purification of cytotoxic granules and cytolytic proteins from NK cells
- · Mouse genetic manipulation by CRISPR-Cas9, and automated genotyping with custom R app

2015 2009

### **Graduate Research Assistant**

Jens Lykke-Andersen's Lab

- University of California San Diego
- · Investigation of co-factors involved in ZFP36-mediated mRNA decay and translational repression during macrophage innate immune response
- · Characterization of the ZFP36 family proteins in cell cycle progression regulation of mouse embryonic fibroblasts

2009

## **Undergraduate Research Assistant**

Ruichuan Chen's Lab

- Xiamen University
- · Exploration of cell cycle and apoptosis implications of the transcription elongation regulator HEXIM1
- · Mechanistic characterization of p-TEFb activation by HIV-Tat



## CONTACT

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github.com/raysinensis

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Bioinformatic tools:

*clustifyr*, automated single cell RNA-seq cell identity assignment Bioconductor package

sauirrelBox. RNA-sea data visualization, exploration, and analysis web browser built with R Shiny and JavaScript

**someta**, quantification and automated monitoring of missing cell-level metadata in scRNA-seq GEO deposition

**scraps**, scalable pipeline for extraction of polyadenylation site and internal priming information from 3' capture scRNA-seq data

Course material:

bioinformatics intro.

rnabioco.github.io/practicaldata-analysis

scRNA-seg workshop. rnabioco.github.io/cellar

2007

	2	PUBLICATIONS, AS FIRST OR CORRESP. AUTHOR
2021		scraps: alternative poly-adenylation site usage pipeline for 3' capture single cell RNA sequencing In Prep for <i>Genome Biology</i> , code at github.com/rnabioco/scraps • Fu R, Riemondy K, Hesselberth J, Jordan C, Gillen A.
2021		Single cell profiling of airway associated CD4+ T cells in chronic beryllium disease Submitted to European Respiratory Journal  · Shaikh A <sup>7</sup> , Fu R <sup>7</sup> , Falta M, Martin A, Yang I, Maier L, Fontenot A.
2021		Transcriptomic response dynamics of human primary and immortalized adrenocortical cells to steroidogenic stimuli <i>Cells</i> · Wellman K <sup>I</sup> , Fu R <sup>I</sup> , Baldwin A, Rege J, Rainey W, Mukherjee N.
2021		RNA-binding proteins regulate aldosterone homeostasis in human steroidogenic cells RNA • Fu R <sup>1</sup> , Wellman K <sup>1</sup> , Baldwin A, Rege J, Bartholomay K, Hirsekorn A, Riemondy K, Rainey W, Mukherjee N.
2021		<b>Cell-level metadata is indispensable for documenting single cell sequencing datasets</b> <i>PLOS Biology</i> • Puntambekar S, Hesselberth J, Riemondy K <sup>c</sup> , <b>Fu R<sup>c</sup></b> .
2021	•	Liver transcriptome dynamics during hibernation are shaped by a shifting balance between transcription and RNA stability  Frontiers in Physiology  Gillen A <sup>1</sup> , Fu R <sup>1</sup> , Riemondy K <sup>1</sup> , Jager J, Fang B, Lazar M, Martin S.
2020		Dynamic RNA regulation in the brain underlies physiological plasticity in a hibernating mammal Frontiers in Physiology • Fu R <sup>7</sup> , Gillen A <sup>7</sup> , Grabek K, Riemondy K, Epperson E, Bustamant C, Hesselberth J, Martin S.
2020		clustifyr: an R package for automated single-cell RNA sequencing cluster classification F1000Research • Fu R, Gillen A, Sheridan R, Tian C, Daya M, Hao Y, Hesselberth J, Riemondy K.
2018	•	PNPT1 release from mitochondria during apoptosis triggers decay of poly(A) RNAs <i>Cell</i> Liu X <sup>7</sup> , Fu R <sup>7</sup> , Pan Y, Meza-Sosa K, Zhang Z, Lieberman J.
2016		Recruitment of the 4EHP-GYF2 cap-binding complex to tetraproline motifs of tristetraprolin promotes repression and degradation of mRNAs with AU-rich elements <i>RNA</i> • Fu R, Olsen M, Webb K, Bennett E, Lykke-Andersen J.
		PUBLICATIONS, AS CONTRIBUTING AUTHOR
2021		Microglia engulf myelin during antibody-mediated cortical demyelination. Submitted to <i>Nature Medicine</i> . Barr H, Given K, McClain C, <b>Fu R</b> , Stockton M, Hickman J, Lee L, Gruber R, Ofengeim D, Macklin W, Bennett J, Owens G, Hughes E.
2021		DUX4 induces the production of truncated RNA binding proteins in human muscle cells. Submitted to <i>eLife</i> , <i>bioRxiv</i> . Campbell A, Dyle M, Calviello L, Matheny T, Cortazar M, Forman T, <b>Fu R</b> , Gillen A, Floor S, Jagannathan S.
2021	•	Post-transcriptional control of mating-type gene expression during gametogenesis in Saccharomyces cerevisiae. <i>Biomolecules</i> . Yeager R, Bushkin G, Singer E, <b>Fu R</b> , Cooperman B, McMurray M.

- Srsf3 mediates alternative RNA splicing downstream of PDGFRa signaling in the facial mesenchyme. Development. Dennison B, Larson E, **Fu R**, Mo J, Fantauzzo K.
- Molecular tracking devices quantify antigen distribution and archiving in the lymph node. *eLife.*Walsh S, Sheridan R, Doan T, Lucas E, Ware B, **Fu R**, Burchill M, Hesselberth J, Tamburini B.
- Single-cell RNA sequencing identifies macrophage transcriptional heterogeneities in granulomatous diseases. *European Respiratory Journal*. Liao S, Shaikh A, Mould K, Konigsberg I, **Fu R**, Davidson E, Li L, Fontenot A, Maier L, Yang I.
- Monocytic Subclones Confer Resistance to Venetoclax-Based Therapy in Acute Myeloid Leukemia Patients. Cancer Discovery. Pei S, Pollyea D, Gustafson A, Stevens B, Minhajuddin M, Fu R, Riemondy K, Gillen A, Sheridan R, Kim J, Costello J, Amaya M, Inguva A, Winters A, Ye H, Krug A, Jones C, Adane B, Khan N, Ponder J, Schowinsky J, Abbott D, Hammes A, Myers J, Ashton J, Nemkov T, D'Alessandro A, Gutman J, Fesik S, Ramsey H, Savona M, Smith C, Jordan C.
- Chronic Liver Disease in Humans Causes Expansion and Differentiation of Liver Lymphatic Endothelial Cells. *Frontiers in Immunology*. Tamburini B, Finlon JM, Gillen A, Kriss M, Riemondy K, **Fu R**, Schuyler R, Hesselberth J, Rosen H, Burchill M.
- Single cell RNA Sequencing Identifies Transforming Growth Factor β as a Critical Regulator of Alveolar Regeneration. *JCI Insight*. Riemondy K, Jansing N, Jiang P, Redente E, Gillen A, **Fu R**, Anthony G, Hesselberth J, Zemans R.
- Pacovery and analysis of transcriptome subsets from pooled single-cell RNA-seq libraries. *Nucleic Acid Research*. Riemondy K, Ransom M, Alderman C, Gillen A, **Fu R**, Finlay-Schultz J, Kirkpatrick G, Di Paola J, Kabos P, Sartorius C, Hesselberth J.

## POSTERS AND PRESENTATIONS

- Practical guide to spatial transcriptomics. (Pulmonary Division Excellence in Respiratory and Critical Care Seminar Series **Invited Presentation**)
- Interactive scRNA-seq cell-type classification identifies widespread omission of cell-level annotations in public data repositories. (Keystone eSymposia on Single Cell Biology Poster with mentee)
- Towards a comprehensive view of dynamic RNA regulation in hibernating 13-lined ground squirrel. (EMBL Conference, From Functional Genomics to Systems Biology Poster)
- ClustifyR: automated single-cell RNA sequencing cluster classification. (Keystone Symposia on Single Cell Biology - Poster)
- PNPT1 release from the mitochondrial intermembrane space triggers rapid decay of mRNA and poly(A)-tailed ncRNA during apoptosis. (Boston Children's Hospital Research Showcase Poster)
- TTP-mediated mRNA repression involves recruitment of the 4EHP-GYF2 complex. (Cold Spring Harbor Laboratory mRNA Processing Meeting **Selected Presentation**)
- TTP family proteins regulate the stability of retinoblastoma protein mRNAs during serum-stimulated G<sub>0</sub>-S transition. (Cold Spring Harbor Laboratory mRNA Processing Meeting Poster)
- The regulation of TTP family mRNA-decay factors during G<sub>0</sub>-S cell cycle progression. (Keystone Symposia on Protein-RNA Interactions Poster)

## ♣☐ TEACHING EXPERIENCE

- Lecture Instructor (Graduate-Level Courses)
- University of Colorado Anschutz
- Foundations in Biomedical Sciences (2018, 2019, 2020 online) lectures, paper discussion, and homework/exam design on RNA biology
- · Workshop for scRNA-seq Data Analysis (2019, 2020 online, 2021) course development and lectures
- · Practical Data Analysis with R/RStudio (2018, 2019) course development and lectures
- $\cdot$  Informatics and Statistics for Molecular Biology (2020 online) teaching and grading on RStudio Cloud
- · Rigor in Research (2020) lecture on RNA sequencing

#### Weekly RNA Bioscience Initiative Informatics Office Hour

University of Colorado Anschutz

· Guidance on RNA and computational research questions from students and researchers of CU Anschutz

#### Teaching Assistant (Graduate-Level Courses)

University of Colorado Anschutz

- · Practical Computational Biology for Biologists: R (2018, 2019)
- · Practical Computational Biology for Biologists: Python (2018, 2019)
- · Core Topics in Biomedical Sciences (2018)

## Teaching Assistant (Undergraduate-Level Courses)

University of California San Diego

- · Molecular Biology (2012)
- · DNA Recombination Lab (2011, 2012)

## **MENTORING EXPERIENCE**

Current 2020

#### **RNA Bioscience Initiative Internship Mentoring**

University of Colorado Anschutz

- · Mentee: AAPI undergraduate computer science student for 2020 summer internship and beyond
- · Project: systemic exploration of published scRNA-seq cell type signatures in the Gene Expression Omnibus
- · Mentee's work was integrated into first-author data-driven essay, updates to R Bioconductor packages clustifyr and clustifyrdatahub, GitHub repo someta, and poster presentation.

2017

#### **Summer Research Mentoring**

Harvard Medical School

- · Mentee: high school student enrolled in the MIT Research Science Institute program
- · Topics: tissue culture, RNA-related bench experiments, RNA-seq informatics analysis, and scientific writing
- · Facilitated research opportunity for mentee at Tsinghua University a year later.

2015 2013

#### BS/MS Student Research Mentoring

University of California San Diego

- · Mentee: female undergraduate/master student
- · Project: biochemical investigation of RNA-binding protein functions
- · Mentee's work was integrated into second-author 2016 RNA publication. Mentee was awarded Best Poster Presentation Award at 2014 UCSD Biological Sciences Annual Student Research Showcase.



## SERVICE AND TRAINING

Current 2019

Annual RNA Bioscience Initiative Grant Review

Current 2018

2016

**GENETICS** Early Career Reviewer (Cellular Genetics section)

16 reviews for Genetics, Science Reports, Frontiers in Immunology, ncRNA, etc; bit.ly/publons-rui-fu

2020

Organizer for CU Anschutz Immuno-Informatics Joint Journal Club

2017 Data Incubator Data Science Fellowship

Harvard Medical School Scientists Teaching Science Course



## DIVERSITY, EQUITY, AND INCLUSION

2021 2019 Admission and Evaluation Committees for CU Graduate Experiences for Multicultural Students Program

2021 RNA Society Volunteer - Writer for Scientist Spotlight 2021 CU Anschutz Medical Campus Equity Certificate Program training and LGBTQ+ Hub Diversity training 2021 **Inclusive STEM Teaching Course training** 2019 Mentor for CU Anschutz and Denver Campus ISCORE undergraduate mentorship program RECOGNITION AND FUNDING 2021 CU Anschutz Medical Campus Spatial Transcriptomics Pilot Grant Award Amount: \$15,000 · Spatiotemporal Analysis of Microglial Contribution to Immune-Mediated Demyelinating Lesion Formation · Formulated plans and drafted proposal, now carrying out experiments and analysis · Joint submission with Dr. Wendy Macklin 2021 Substantial NIH Uo1 Grant Writing Submitted to National Institute of Diabetes and Digestive and Kidney Diseases (NIDDK) · Spatial Mapping of Proteomic and Transcriptional Signatures in Kidney Disease · Drafted all content and preliminary data on spatial RNA profiling (~25% of entire proposal) · In collaboration with Dr. Joshua Thurman, Dr. Elena Hsieh, and Dr. Julia Wrobel 2021 Colorado RNA Newsletter Spotlight 2011 Dr. Huang Memorial Scholarship, UC San Diego Award Amount: \$108,000 2009 2008 First-Class Scholarship, XMU University Merit Student 2006 SCIENCE OUTREACH

- Facilitator at Denver Museum of Nature and Science Prehistoric Journey section (2019 2020)
- Visitor Education Volunteer at the New England Aquarium (2016 2017)
- Information Ambassador at the San Diego Zoo (2015 2016)
- Organizer for UCSD Ethics Center's Silent Spring 50th Anniversary Project (2012)

## **REFERENCES**

## Dr. Jens Lykke-Andersen, Professor

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### Dr. Judy Lieberman, Professor and Chair

Cellular and Molecular Medicine, Boston Children's Hospital of Harvard Medical School

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#### Dr. Jay Hesselberth, Associate Professor

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Code at github.com/raysinensis/cv.

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