# Joha Park, Ph.D.

#### **KEY SKILLS & EXPERIENCE**

- Computational and Molecular RNA Biology
- Sequencing Data Analysis: (sc)RNA-seq, CLIP-seq, etc.
- Tissue Clearing and Expansion for Spatial Genomics
- Imaging Data Analysis: 3D Confocal/Light-sheet
- Multi-omics, Image Processing, Machine Learning

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Institute of

Technology

#### **CURRENT POSITION**

## 01/2020 - present Postdoctoral Associate/Fellow

Chung Lab (webpage) (Advisor: Kwanghun Chung)
Institute for Medical Engineering and Science
The Picower Institute for Learning and Memory
Massachusetts Institute of Technology, Cambridge, MA, US

## Roles

- Developing a new method for hydrogel-based tissue engineering
- Building a unified and scalable analytical framework for 3D light-sheet microscopy
- Generation of a single-cell mouse brain reference atlas

#### **PROFESSIONAL EXPERIENCE**

## 2019 Postdoctoral Fellow

Center for RNA Research (Director: V. Narry Kim) Institute for Basic Science Seoul National University, Seoul, Korea

#### Roles

- Leading projects on mRNA poly(A)-tail length regulations
- Conventional and custom NGS library prep & data analysis
- Wet and dry experiments related to cell and RNA biology

## **EDUCATION AND TRAINING**

2012 – 2019 **Ph.D. in Biological Sciences** (RNA Biology & Bioinformatics)

RNA Biology Lab (Advisor: V. Narry Kim)

Dissertation: "Poly(A) length regulation: deadenylases and

the poly(A) barricade"

2015 - 2018 Technical Research Personnel

**Compulsory Military Service** 

2008 – 2012 B.S. in Biological Sciences (cum laude)

Seoul National University

Seoul National University

ROK Army

Seoul National University

### **PUBLICATIONS**

\*: co-first author, #: co-corresponding author

1 J. Park\*, M. Kim\*, H. Yi\*, K. Baeg\*, Y. Choi, Y.-s. Lee, J. Lim, V. N. Kim (2022) "Short poly(A) tails are protected from deadenylation by the LARP1-PABP complex" Nat. Struct. Mol. Biol. (accepted)

- 2 J. Park\*, S. Khan\*, D. H. Yun\*, T. Ku, K. L. Villa, J. E. Lee, Q. Zhang, G. Feng, J. Park, E. Nedivi#, K. Chung# (2021) "Epitope-preserving magnified analysis of proteome (eMAP)" Sci. Adv. 7(16):eabf6589
- 3 Y. Kim\*#, <u>J. Park\*</u>, S. Kim\*, M. Kim, M.-G. Kang, C. Kwak, M. Kang, B. Kim, H.-W. Rhee, and V. N. Kim# (2018) "PKR senses nuclear and mitochondrial signals by interacting with endogenous double-stranded RNAs" *Mol. Cell*, 71(6):1051–1063.e6
  - Selected as "Research Highlights" in Nat. Chem. Biol., 14(11):989
- 4 H. Yi\*, <u>J. Park\*</u>, M. Ha, J. Lim, H. Chang and V. N. Kim (2018) "PABP cooperates with the CCR4-NOT complex to promote mRNA deadenylation and block precocious decay" *Mol. Cell*, 70(6):1081–1088.e5
  - Previewed in *Mol. Cell*, 70(6):987–988
- 5 T. A. Nguyen\*#, <u>J. Park\*</u>, T. L. Dang, Y.-G Choi, and V. N. Kim# (2018) "Microprocessor depends on hemin to recognize the apical loop of primary microRNA" *Nucleic Acids Res.*, 46(11):5726–5736
- 6 K. T. You, **J. Park**, and V. N. Kim (2015) "Role of the small subunit processome in the maintenance of pluripotent stem cells" *Genes Dev.*, 29(19):2004–2009
- 7 T. A. Nguyen, M. H. Jo, Y.-G. Choi, **J. Park**, S. C. Kwon, S. Hohng, V. N. Kim# and J.-S. Woo# (2015) "Functional anatomy of the human Microprocessor" *Cell*, 161(6):1374–1387
- 8 Y.-K. Kim, G. Wee, **J. Park**, J. Kim, D. Baek, J.-S. Kim#, and V. N. Kim# (2013) "TALENbased knockout library for human microRNAs" *Nat. Struct. Mol. Biol.*, 20(12):1458–1464

Links: Google Scholar, PubMed

## **HONORS AND AWARDS**

- 04/2022 present Picower Postdoctoral Fellows, The Picower Institute for Learning and Memory, MIT
  - 2021 AKN Outstanding Research Awards, Association of Korean Neuroscientists
  - 2019 Conference Scholarship, Keystone Symposia: Small Regulatory RNAs
  - 2019 Best Dissertation Awards, School of Biological Sciences, SNU
  - 2008 2012 Presidential Science Scholarship, Korea Science and Engineering Foundation

# INTERNATIONAL CONFERENCES

- 2019 Cell Symposia: Regulatory RNAs *short talk*"Poly(A) barricade as a rate-limiting factor that shapes poly(A) tail length"
- 2018 CSHL Meeting: Regulatory & Noncoding RNAs *short talk*"PKR senses nuclear and mitochondrial signals by interacting with endogenous dsRNAs"
- 2017 CSHL Meeting: Eukaryotic mRNA Processing *short talk* "Genome-wide evaluation of the role and specificity of deadenylases"
- 2016 EMBL Symposium: The Complex Life of mRNA poster
- 2016 IMBA: 11th Microsymposium on Small RNAs poster
- 2014 Keystone Symposia: RNA Silencing poster

### **INVITED TALKS**

05/2022 Special seminar – *Department of Biomedical Engineering, UNIST*"Hydrogel-based tissue engineering for holistic interrogation of biological systems"

### **TEACHING EXPERIENCE**

- 2014 Teaching Assistant, Advanced RNA Biology, School of Biological Sciences, SNU
- 2013 Teaching Assistant, Biology Lab 2, School of Biological Sciences, SNU
- 2012 Teaching Assistant, Biology Lab 1, School of Biological Sciences, SNU
- 2011 Tutor, Basic Course for Biology, School of Biological Sciences, SNU

### **TECHNICAL EXPERTISE**

## **Computational biology**

- General skills
  - Unix/Linux environment
  - Programming languages: Python, R, Julia
  - Building a custom analysis pipeline: Snakemake, GNU Make
  - Efficient data visualization: Matplotlib, Seaborn, Plotly
- RNA biology
  - (sc)RNA-seq, Ribo-seq, CLIP-seq, ATAC-seq, TAIL-seq, etc.
  - Statistical analysis for genome-wide, transcriptomic studies
  - Tools: Pandas, Scanpy, Squidpy, Seurat, STAR, Bedtools, Samtools, etc.
- Image processing
  - Concurrent programming for large-scale images
  - 2D/3D rigid and non-rigid biomedical image registration
  - Convolutional Neural Networks
  - Tools: Numpy, Scipy, PyTorch, Scikit-learn/image, OpenCV, ImageJ/Fiji, etc.

## **Wet experiments**

- Routine experiment
  - Molecular cloning: Gibson assembly, Golden Gate assembly
  - Mammalian & bacterial cell culture
  - DNA and siRNA transfection
  - Quantitative RT-PCR
  - Immunoprecipitation
  - Western blotting
- RNA biology
  - Custom RNA sequencing library preparation
  - Custom high-resolution poly(A)-tail length assay
  - Tet-On inducible reporter cell line generation
  - RNA co-immunoprecipitation
  - Lentiviral transduction
  - In vitro transcription
  - Northern blotting
  - Luciferase reporter assay