

BIOGRAPHICAL SKETCH

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NAME: Hong Jin

eRA COMMONS USER NAME (credential, e.g., agency login): HongJin

POSITION TITLE: Assistant Professor of Biochemistry

EDUCATION/TRAINING *(Begin with baccalaureate or other initial professional education, such as nursing, include postdoctoral training and residency training if applicable. Add/delete rows as necessary.)*

| INSTITUTION AND LOCATION | DEGREE (if applicable) | Completion Date MM/YYYY | FIELD OF STUDY |
|--|---------------------------|-------------------------------|--------------------------------------|
| Central China Normal University, Wuhan, Hubei, China | B.S. | 05/1995 | Chemistry |
| Wuhan University, Wuhan, Hubei, China | M. S. | 05/1998 | Analytical Chemistry |
| University of Massachusetts at Boston, Boston, MA | | 05/2000 | Physical Chemistry |
| Yale University, New Haven, CT, USA | Ph.D. | 05/2007 | Biophysics |
| Medical Research Council (MRC) - Laboratory of Molecular biology, Cambridge, UK | Postdoctoral training | 01/2008- 12/2012 | Biophysics, X-ray Crystallography |

A. Personal Statement

My research focuses on understanding the molecular mechanisms of translation and translational regulation in the cell. My group is best known for revealing structures and functions of the ribosome, as well as RNAs and RNA-binding proteins involved in the cellular translation process that is fundamental to gene expression and regulation. I have broad expertise in Biochemistry, Biophysics and Structural Biology including macromolecule X-ray crystallography, solution NMR spectroscopy and single particle cryoEM. As a Ruth L. Kirschstein National Research Postdoctoral Fellow, I carried out extensive biochemical investigations in translation and solved several high-resolution crystal structures of ribosomal complexes with protein release factors bound along the translational termination pathway by X-ray crystallography. These include the first high-resolution crystal structure of a translational GTPase, the class II release factor RF3, bound to the hybrid state of the ribosome. These structures have addressed fundamental questions in molecular biology that have persisted since the genetic code was discovered by revealing universally conserved protein-RNA interactions that contribute to the specificity of stop-codon recognition and the catalysis of peptide release on the ribosome. I also characterized the molecular features of the ratcheting ribosome and demonstrated how a GTPase induces and stabilizes the ribosome in the hybrid state. These features are fundamental to the ribosome function in each of its essential stages of translation including initiation, elongation, termination and recycling. As a Ph.D student at Yale University, I initiated a biochemical investigation of eukaryotic small nucleolar RNA (snoRNA) and ribosomal RNA (rRNA) interactions, and solved the first structure of a snoRNA-rRNA complex by solution NMR spectroscopy. My dissertation work revealed a new RNA-RNA interaction motif, named as the Ω motif, which provided mechanistic insights into how a specific nucleotide in the rRNA is selected for modifications that are essential for cellular functions. As a young investigator, I have laid the groundwork for our research by not only developing effective experimental strategies and protocols, but also providing leadership and administrative skills. Our key findings were published in the peer-reviewed journals including *Nature*, *PNAS*, *RNA* and *Scientific Reports*. I believe my group is uniquely positioned to investigate the fundamental questions that require a combination of calculated risk-taking and a focused desire to tackle important yet underexplored areas of science. My scientific training, experience and past success in RNA and translation will certainly be of great value in pioneering our research and I look forward to the challenges and opportunities that confront our investigations.

B. Positions and Honors

Research and Professional Positions

| | |
|-------------------|--|
| 01/2008 – 12/2012 | Postdoctoral Fellow (Supervisor: Dr. Venki Ramakrishnan) MRC Laboratory of Molecular Biology, Cambridge, UK |
| 09/2009 – 12/2012 | Research Fellow in Science (Equivalent to a Faculty Position in the US) University of Cambridge, Lucy Cavendish College, Cambridge, UK |
| 12/2012 – Present | Assistant Professor of Biochemistry Department of Biochemistry, Center for Biophysics and Quantitative Biology University of Illinois at Urbana-Champaign |

Academic and Professional Honors

| | |
|-----------|---|
| 2009-2012 | NIH Ruth L. Kirschstein National Research Service Award, USA <ul style="list-style-type: none">The award includes my postdoctoral salary and institutional allowance that covers my conference travel and part of the research expense. |
| 2009-2012 | Ethel Cruickshank Research Fellowship, University of Cambridge - Lucy Cavendish College, UK <ul style="list-style-type: none">The award is equivalent to a faculty position in the US, which includes participating and providing various academic services such as seminars, lectures, governing body meetings and other administrative services. It also includes lively participation in the college life and activities in the Lucy Cavendish College and other colleges in the University of Cambridge. |
| 2002 | Yale University Extraordinary Teaching Assistant Award <ul style="list-style-type: none">The award is based upon student evaluations and nominations |
| 1992-1995 | Central China Normal University Scholarship for Academic Excellence <ul style="list-style-type: none">Top 2% of the university |

Memberships in professional societies:

| | |
|--------------|------------------------------------|
| 2008-Present | The RNA Society, USA |
| 2009-Present | The Biophysical Society, USA |
| 2007-Present | The American Chemical Society, USA |

Other Experience and Professional Activity

| | |
|---------------|--|
| 2007- Present | Invited speaker at 16 institutions and conferences |
| 2016 | Ad Hoc reviewer for NSF (Systems and Synthetic Biology) |
| 2016 | Ad Hoc reviewer for NIH as an early stage investigator (GM-MSFC) |

C. Contribution to Science

1. An induced-fit mechanism for co-translational quality control process in the ribosome

Intervening protein synthesis when an error occurs is an essential quality control mechanism that contributes to the overall accuracy and fidelity of translation in living cells. In this process, cellular signals that initiate quality control process are first deciphered in the translating ribosome, and then release factors or release factor-like proteins are recruited to terminate protein synthesis, followed by recycling of ribosomal subunits and degradation of faulty translational components. Using ArfA and RF2 as a model system, we determined the kinetics of ArfA/RF2-mediated co-translational quality control on bacterial ribosome follows an induced-fit mechanism. Since the signals that intervention is required originate in the decoding center of the small ribosomal subunit while the actions that result in termination of protein synthesis occur in the large subunit of the ribosome, *the induced-fit mechanism for co-translational quality control that we have proposed is likely to be universally conserved.*

Our results have a broader impact towards uncovering new functions of the ribosome in quality control pathways, demonstrating that beyond making proteins in living cells, ribosomes actively maintain translation accuracy and fidelity by recruiting proteins other than the canonical translational factors.

My role in this work is the PI. I designed experiments, and performed most of the experiments with a postdoctoral fellow in my laboratory in this research.

- Fuxing Zeng and **Hong Jin**
Peptide release promoted by methylated RF2 and ArfA in nonstop translation is achieved by an induced-fit mechanism *RNA*, 22(1): 49–60, 2016 PMID: PMC4691834
- Fuxing Zeng, Yanbo Chen, Jonathan Remis, Mrinal Shekhar, James C. Phillips, Emad Tajkhorshid and **Hong Jin**.
Structural basis of co-translational quality control by ArfA and RF2 bound to ribosome. *Nature*, 541(7638), 554-557, 2017 PMID: 28077875

2. A dual role of the decapping activator in the eukaryotic cell.

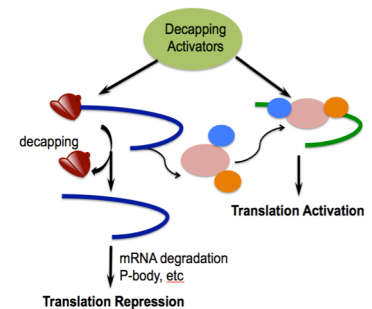
Decapping activators facilitate removal of the 5' m⁷G caps from eukaryotic mRNAs by assisting the assembly of decapping complexes or stimulating activities of the decapping enzyme, and they are known to be translational repressors to inhibit translation and promote storage or degradation of mRNAs in cytoplasm. Using biochemistry, we showed that one decapping enhancer, Sbp1, selectively promotes the translation of mRNA encoding the polyadenosine-binding protein (Pab1) and other mRNAs possessing cap-independent translation initiation activities. We further demonstrated molecular interactions important for Sbp1-specific translational regulation and the underlying molecular mechanism.

Our results not only reveal a dual role of the decapping activator in regulating mRNA translation: a general translation repressor and a translation activator for subset of mRNAs in the cell, but also connect the two seemingly unrelated processes, decapping and translation activation of cellular mRNAs, thereby identifying a new layer of translational control in eukaryotic cells.

My role in this work is PI. I assembled a strong research team, designed experiments and participated the experimental work with my team.

- Alberto Brandariz-Núñez, Fuxing Zeng, Quan Ngoc Lam and **Hong Jin**.

Sbp1 modulates the translation of Pab1 mRNA in a poly(A)- and RGG-dependent manner, *RNA*, 24, 43-55, 2018 PMID: 28986506

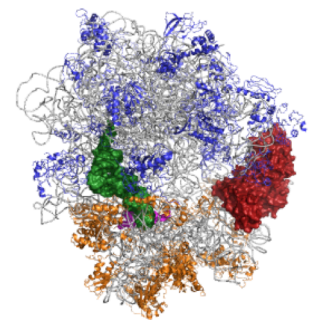


3. Solve the first high-resolution crystal structure of a translational GTPase bound to the hybrid state of the ribosome by X-ray crystallography

Translational GTPases ensure the speed, fidelity and accuracy in each step of the translation. Ribosomes recruit the translational GTPases in the hybrid ligand-binding states where the two ribosomal subunits rotate relative to one another. Following a thorough biochemical characterization of the translation system, I solved the first high-resolution crystal structure of a translational GTPase, the class II release factor 3 (RF3), bound to the hybrid state of the ribosome by X-ray crystallography. The structure revealed the molecular interactions on how a GTPase induces and stabilizes the hybrid state of the ribosome and shed light on the function of RF3 in translational termination. My work also elucidated the structural features of the ratcheting ribosome in the hybrid ligand-binding state that are the fundamental feature of the ribosome function in the essential stages of translation including initiation, translocation, termination and recycling.

Furthermore, the new crystal form that I discovered has paved the way for solving new structures of ribosomes with other translational GTPases bound. These structures have opened a new field for functional investigations and have provided enormous molecular details and framework for designing new antibiotics targeting bacterial ribosome functions.

I led this project as a postdoctoral fellow. I designed and performed all the experimental work while one technician in the lab helped on a protein purification step.



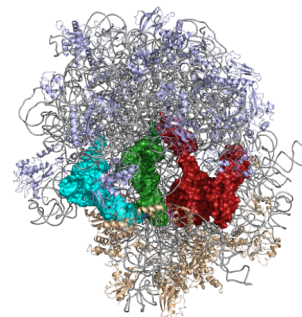
- **Hong Jin**, Ann C. Kelley and V. Ramakrishnan
Crystal structure of the hybrid state of ribosome in complex with the GTPase release factor 3.

4. Solve the first high-resolution crystal structure of a protein release factor bound to the 70S ribosome

The specificity of stop-codon recognition and the catalytic mechanism of peptide release have been the fundamental questions in molecular biology since the elucidation of the genetic code. I achieved the major breakthrough in structural studies on ribosomal complexes in the first year of my postdoctoral training with Dr. Venki Ramakrishnan. I established a new and reproducible condition for obtaining high-resolution 70S ribosome crystals with protein factors bound and *determined the first high-resolution crystal structure of release factor 2 (RF2) bound to the 70S ribosome*. The structure revealed universally conserved protein-RNA interactions in the ribosome during termination. I also obtained structures of other ribosomal complexes along the termination pathway. Together, these structures answered long-standing questions about the specificity of stop-codon recognition and addressed the catalytic mechanism of peptide release.

Furthermore, the condition that I have established for 70S ribosome crystallization and cryo-protection has paved way for the subsequent success on obtaining structures of other bacterial ribosomal complexes in the Ramakrishnan laboratory.

I led this project to the successful completion as a postdoctoral fellow. I designed the sample preparation and crystallization plan. I performed the experiments and screened the crystals with help from my colleagues.



- Albert Weixlbaumer*, **Hong Jin***, Cajetan Neubauer, Rebecca M. Voorhees, Sabine Petry, Ann C. Kelley and V. Ramakrishnan (****These two authors contributed equally to the work***)
Insights into translational termination from the structure of RF2 bound to the ribosome.
Science, 322(5903): 953-956, 2008 PMCID: PMC2642913
- **Hong Jin**, Ann C. Kelley, David Loakes, and V. Ramakrishnan
The structure of the 70S ribosome bound to RF2 and a substrate analog provides insights into catalysis of peptide release.
Proceedings of the National Academy of Sciences, 107(19): 8593-8598, 2010
PMCID: PMC2889298

5. Solve the first structure of the human U65 snoRNA bound to rRNA by solution NMR spectroscopy

Noncoding RNAs play essential and diverse roles in regulating gene expression. My Ph.D. work focused on one type of the noncoding RNAs in eukaryotic cells, small nucleolar RNAs (snoRNAs). Base pairings between snoRNAs and sequences in eukaryotic rRNAs target specific nucleotides for modifications. Most of the modifications are clustered in the functionally important regions of the ribosome. rRNA modifications are critical for ribosome biogenesis, assembly, structure and translating function. snoRNAs play other essential functions in the cell, including participating in pre-mRNA processing, directing alternative splicing and acting as microRNA precursors. As a Ph.D. student, I developed a model system, used extensive biochemical and biophysical methods including chemical and enzymatic footprinting, native gel binding assay, electrospray mass spectrometry, and analytical ultracentrifugation experiment, to characterize the model and parent systems, and subsequently *determined the first structure of the human U65 snoRNA with and without substrate bound by solution NMR spectroscopy*. This structure revealed a new Ω -shaped RNA interaction motif that is conserved in all box H/ACA snoRNP-substrate complexes and provided the first physical evidence to support unique features of the substrate-recognition by box H/ACA snoRNA.



My role in this work is a Ph.D. student. I designed and carried out all the experimental investigations in this research. My Ph.D. research provided *the first structural basis for box H/ACA snoRNA-mediated pseudouridylation in eukaryotic cells*.

- **Hong Jin**, J. Patrick Loria and Peter B. Moore.
Solution structure of an rRNA substrate bound to the pseudouridylation pocket of a box H/ACA snoRNA.
Molecular Cell, 26(2): 205-215, 2007 PMID: 17466623

Other publications in Chemistry and Analytical Chemistry fields are:

- Fuxing Zeng and **Hong Jin**.
Conformation of methylated GGQ in the Peptidyl Transferase Center during Translation Termination.
Scientific Reports, 8, 2349, 2018. PMID: 29403017
- Hong Wang, Yuan-yuan Zhao, **Hong Jin**, and Hua-shan Zhang
N-hydroxy-succinimidyl- α -naphthylacetate as a derivatizing reagent for amino acids and oligopeptides in RP-HPLC.
Journal of Liquid Chromatography, 24(20): 3157-3170. 2001
- Hong Wang, **Hong Jin** and Hua-shan Zhang
Determination of catecholamines as their N-hydroxy-succinimidyl-3-indolylacetate derivatives by pre-column derivatization HPLC separation and fluorescent detection.
Fresenius Journal of Analytical Chemistry, 365(8): 682-684. 1999
- **Hong Jin**, Xun Liu, Hong Wang, Hua-shan Zhang and Jie-ke Cheng
Determination of amino acids as their N-hydroxy-succinimidyl-3-indolylacetate derivatives by pre-column derivatization HPLC separation and fluorescent detection.
Wuhan Da Xue Xue Bao (Natural Science Journal of Wuhan University, China), 44(2): 175-178, 1998
- Guanghan Lu, **Hong Jin** and Dandan Song
Determination of trace nitrite by anodic stripping voltammetry.
Food Chemistry, 59(4): 583-587. 1997

List of Published Work in MyBibliography:

http://www.ncbi.nlm.nih.gov/sites/myncbi/18WK9L8_bYGAJ/bibliography/48229889/public/?sort=date&direction=descending

D. Research Support

Ongoing Research Support:

NIH R01GM120552
Title: Mechanisms of Translational Activation by Decapping Activators
Role: Hong Jin, **P.I.**
Award period: 9/22/16 – 08/31/2021

Completed Research Support:

NIH F32GM087083 NIH Ruth L. Kirschstein National Research Service Award
Title: Characterization of Class I Release Factor-Mediated Translation Termination
Role: Hong Jin, **P.I.**
Award period: 2009-2012