Welcome to the Department of Biochemistry and Molecular Biophysics

Washington University in St. Louis
School of Medicine

https://biochem.wustl.edu
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2) Click **Media and Newsletters**

3) Click **Display TV Announcements**

View these slides online!
March Publication

Min Kyung Shinn, Sumit K. Chaturvedi, Alexander G. Kozlov, & Timothy M. Lohman

Allosteric effects of E. coli SSB and RecR proteins on RecO protein binding to DNA

The **Burgers Lab** studies DNA replication and DNA damage response in eukaryotic cells. Using yeast as a model organism, the lab integrates the biochemical analysis of DNA-protein interactions in purified model systems with the genetic analysis of targeted yeast mutants. Specific areas of interest are lagging strand DNA replication and Okazaki fragment maturation, damage induced mutagenesis, and DNA damage cell cycle checkpoints.

Right: DNA replication fork and Okazaki fragment maturation

See more research: [biochem.wustl.edu/spotlight](http://biochem.wustl.edu/spotlight)
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Oct 1st 2022 - Alex Holehouse, assistant professor of biochemistry and molecular has received more than $450,000 in funding from the National Science Foundation (NSF) to lead a five-year project titled “Molecular engineering to understand desiccation protection and water responsiveness.” The project is part of a larger grant called “Life without water: protecting macromolecules, cells, and organisms during desiccation and rehydration across kingdoms of life.” This grant establishes the Water and Life Interface Institute led by Carnegie Science. The new initiative includes collaborators from at least nine research institutions nationwide.
“Molecular-scale mapping of the Pseudomonas aeruginosa biofilm matrix”

Shana Elbaum, Ph.D.
CUNY Advanced Science Research Center

Tuesday, May 23rd, 2023
10:30 am
Biochemistry Seminar Room, 264 McDonnell Science
The **Galburt Lab** strives to understand the physical mechanisms of transcription initiation and other important DNA-protein interactions. More specifically, we use a variety of single-molecule and ensemble biophysical techniques including both optical and magnetic tweezers and fluorescent microscopy to investigate how the assembly of initiation complexes on gene promoters leads to DNA unwinding and transcription. Our work is currently focused on the mechanisms of basal transcription initiation in Eukaryotes and on factor-regulated transcription in Mycobacterium tuberculosis.

See more research: [biochem.wustl.edu/spotlight](http://biochem.wustl.edu/spotlight)
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The **Niemi Lab** investigates how mitochondria are built, regulated, and maintained across physiological contexts. We blend biochemistry, systems biology, and physiology to understand mechanisms of mitochondrial regulation and how they influence metabolism and organellar function. Using insights gained from our molecular studies, we aim to understand how mitochondrial dysfunction contributes to mammalian pathophysiology, with the long-term goal of translating our discoveries into new therapeutic options to restore mitochondrial function in human disease.

See more research: [biochem.wustl.edu/spotlight](http://biochem.wustl.edu/spotlight)
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Garrett M. Ginell & Alex S. Holehouse

An Introduction to the Stickers-and-Spacers Framework as Applied to Biomolecular Condensates

The **Cooper Lab** is interested in how the actin filaments in cells assemble and how that assembly controls cell shape and movement. One focus is an actin-binding protein called "capping protein," which caps one end of the actin filament. Capping protein is in turn regulated by intrinsically disordered regions of the CARMIL family of proteins, which exhibit positive linkage in their binding interactions.

See more research: [biochem.wustl.edu/spotlight](http://biochem.wustl.edu/spotlight)
Research in the **Lohman Lab** focuses on obtaining a molecular understanding of the mechanisms of protein-nucleic acid interactions involved in DNA metabolism, in particular, DNA motor proteins (helicases/translocases) and single stranded DNA binding proteins. Thermodynamic, kinetic, structural and single molecule approaches are used to probe these interactions at the molecular level.

See more research: [biochem.wustl.edu/spotlight](http://biochem.wustl.edu/spotlight)
Are you paid monthly?

Please remember that your time report is due by the 5th of each month.
HAVING ISSUES AT WORK?
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Jessica Kennedy – Title IX Director, jwkennedy@wustl.edu, 314-935-3118
Jessica Kuchta-Miller – Staff/Postdoc/Graduate Student Ombuds, 314-379-8110
Karen O’Malley – Medical Student Ombuds, 314-660-2089
Jim Fehr – Faculty Ombuds, 314-660-2089
Oct 1st 2022 – **Alex Holehouse**, assistant professor of biochemistry and molecular has received $380,000 in funding from the Human Frontiers Science Program (HFSP) to lead a three-year project titled **“Molecular determinants of evolutionary conservation in disordered protein regions”**. This project will integrate computational and experimental approaches to uncover who intrinsically disordered protein regions evolve. The proposal involves co-investigators Dr. Hyun Kate Lee (University of Toronto, CA) and Dr. Dolf Weijers (Wageningen University, NL).
Congratulations to Anna Damato for being selected as the 2022 Ceil M. DeGutis Prize Fellow

Anna Damato is a fifth-year PhD candidate in Neuroscience in the Department of Biology. Anna was nominated for this award by her thesis mentor, Dr. Erik Herzog, in whose lab she is connecting the bench to the bedside by investigating mechanisms of glioblastoma brain tumor circadian rhythms and how they impact the efficacy of chemotherapy. Anna uses real-time bioluminescence reporters of circadian gene expression to analyze the effects of timed treatment, with the goal of maximizing anti-tumor effects and minimizing side effects of chemotherapy in treating an otherwise dismal disease.

Visit biochem.wustl.edu/news to read more!
The **Bowman Lab** seeks to understand the distribution of different structures a protein adopts and how this ensemble determines a protein’s function. Examples of ongoing research projects include 1) understanding how mutations in the enzyme beta-lactamase change its specificity without changing the protein’s crystal structure, 2) designing allosteric drugs, and 3) developing algorithms for quickly building models of the different structures a protein adopts.

See more research: [biochem.wustl.edu/spotlight](http://biochem.wustl.edu/spotlight)
The tetraspanin CD53 protects stressed hematopoietic stem cells via promotion of DREAM complex-mediated quiescence

<table>
<thead>
<tr>
<th>Holiday</th>
<th>Day Observed</th>
<th>Date Observed at WashU</th>
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<tbody>
<tr>
<td>New Year's Eve</td>
<td>Friday</td>
<td>December 30th, 2022</td>
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<td>New Year's Day</td>
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<td>Martin Luther King, Jr.</td>
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<td>January 16th, 2023</td>
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<td>Labor Day</td>
<td>Monday</td>
<td>September 4th, 2023</td>
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The **Greenberg Lab** focuses on how cytoskeletal motors function in both health and disease. Currently, the lab is studying mutations that cause familial cardiomyopathies, the leading cause of sudden cardiac death in people under 30 years old. The lab uses an array of biochemical, biophysical, and cell biological techniques to decipher how these mutations affect heart contraction from the level of single molecules to the level of engineered tissues. Insights into the disease pathogenesis will guide efforts to develop novel therapies.

See more research: [biochem.wustl.edu/spotlight](http://biochem.wustl.edu/spotlight)
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