

Yu Zhang, Ph.D.

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Postdoctoral Research Fellow

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Wyss Institute (Harvard), MIT, and Broad Institute

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Focus: AI-driven, safety-aware antibiotic discovery using deep learning.

I am a Wyss–MIT–Broad postdoc leading cross-disciplinary teams to design generative and multitask models that discover non-toxic antibiotics against Gram-negative pathogens.

Metrics Snapshot: 18 peer-reviewed publications (7 first-author); h-index: 12; >360 citations. Flagship works in *Cell*, *PNAS*, *Nexus*, *iScience*, *npj Systems Biology and Applications*. Competitive recognition includes a departmental top-student award and national society travel support.

Key Independent Contributions:

- **Safety-aware antibiotic discovery:** Established an independent research line that couples multitask toxicity modeling with AI-guided screening to enable discovery of non-toxic antibiotics (under review; senior author team: Collins lab).
- **Systems pharmacology of vascular and immune therapies:** Led mechanistic and QSP studies that uncovered druggable control points in angiogenesis and immuno-oncology, resulting in multiple first-author papers and a PhD thesis.
- **Translational modeling across sectors:** Demonstrated the utility of systems pharmacology in drug development by delivering QSP models used at CytomX Therapeutics to inform dose selection in immuno-oncology trials.

Research Experience

Postdoctoral Research Fellow

2024 – Present

Wyss Institute (Harvard), MIT, and Broad Institute

Cambridge, MA

- **Multitask Learning Architect:** Design and lead a deep learning platform for high-fidelity antibiotic toxicity prediction, enabling discovery of non-toxic clinical candidates (under review).
- **Generative AI Platform:** Contributed to the development and validation of a deep learning framework for *de novo* antibiotic design, with experimentally validated hits against resistant pathogens.
- **Cross-Institutional Leadership:** Direct collaborations across Harvard/Wyss, MIT, and Broad to align *in silico* predictions with high-throughput wet-lab validation.

Ph.D. Candidate

2017 – 2024

Popel Systems Biology Laboratory, Johns Hopkins University

Baltimore, MD

- **Mechanistic Modeling Lead:** Conceived and led the development of mechanistic ODE models of VEGF, Ang-Tie, and integrin signaling to reveal non-intuitive therapeutic targets for vascular stability.
- **QSP Platform Development:** Built and deployed multi-scale biophysical platforms that predict drug efficacy and guide dosing strategies for sickle cell anemia and ischemic diseases.

Industry QSP Intern

Summer 2022

CytomX Therapeutics

South San Francisco, CA

- **Clinical Translation:** Developed clinical QSP models for Probody™ therapeutics used to support dose selection and define safety margins in immuno-oncology trials.

Education

Ph.D. in Biomedical Engineering (GPA: 4.0/4.0)

2017 – 2024

Johns Hopkins University School of Medicine

Baltimore, MD

B.S. in Biomedical Engineering & Applied Mathematics

2013 – 2017

Johns Hopkins University

Baltimore, MD

(Minor: Spanish, GPA: 3.96/4.0; General & Departmental Honors)

Selected Publications & Manuscripts

A generative deep learning approach to de novo antibiotic design. *Cell* (2025)

Krishnan A*, Anahtar MN*, Valeri JA*, ... Zhang Y ... and Collins JJ.

Impact: First demonstration of generative deep learning yielding experimentally validated antibiotic candidates.

Multitask learning enables discovery of nontoxic antibiotics. (under review, 2025)

Zhang Y*, Hennes A*, Krishnan A* ... and Collins JJ.

Impact: Establishes a multitask model for toxicity prediction to overcome a primary bottleneck in drug discovery.

Signaling-biophysical modeling unravels mechanistic control of RBC phagocytosis and clearance in sickle cell disease. *PNAS Nexus* (2024)

Zhang Y, Qiang Y, Li H, ... and Zhao C.

Impact: Established a multi-scale framework linking biophysical mechanics to systemic anemia in sickle cell disease.

Promoting vascular stability through Src inhibition and Tie2 activation. *iScience* (2025)

Zhang Y, Kontos CD, Annex BH, and Popel AS.

Impact: Model-driven identification of combination therapies for vascular normalization.

* indicates co-first authors.

Full Publication List

Peer-reviewed articles

- [1] Aarti Krishnan, Melis N Anahtar, Jacqueline A Valeri, Wengong Jin, Nina M Donghia, Leif Sieben, Andreas Lutten, Yu Zhang, Seyed Majed Modaresi, Andrew Hennes, et al. A generative deep learning approach to de novo antibiotic design. *Cell* (2025).
- [2] Yu Zhang, Christopher D Kontos, Brian H Annex, and Aleksander S Popel. Promoting vascular stability through Src inhibition and Tie2 activation: A model-based analysis. *iScience* **28.6** (2025).
- [3] Alberto Ippolito, Hanwen Wang, Yu Zhang, Vahideh Vakil, Hojjat Bazzazi, and Aleksander S Popel. Eliciting the antitumor immune response with a conditionally activated PD-L1 targeting antibody analyzed with a quantitative systems pharmacology model. *CPT: Pharmacometrics & Systems Pharmacology* **13.1** (2024), pp. 93–105.
- [4] Alberto Ippolito, Hanwen Wang, Yu Zhang, Vahideh Vakil, and Aleksander S Popel. Virtual clinical trials via a QSP immuno-oncology model to simulate the response to a conditionally activated PD-L1 targeting antibody in NSCLC. *Journal of Pharmacokinetics and Pharmacodynamics* **51.6** (2024), pp. 747–757.
- [5] Geli Li, Yuchen Ma, Sujie Zhang, Wen Lin, Xinyi Yao, Yating Zhou, Yanyong Zhao, Qi Rao, Yuchen Qu, Yuan Gao, et al. A mechanistic systems biology model of brain microvascular endothelial cell signaling reveals dynamic pathway-based therapeutic targets for brain ischemia. *Redox Biology* **78** (2024), p. 103415.
- [6] Yu Zhang, Yuhao Qiang, He Li, Guansheng Li, Lu Lu, Ming Dao, George E Karniadakis, Aleksander S Popel, and Chen Zhao. Signaling-biophysical modeling unravels mechanistic control of red blood cell phagocytosis by macrophages in sickle cell disease. *PNAS nexus* **3.2** (2024), pgae031.
- [7] Samira Anbari, Hanwen Wang, Yu Zhang, Jun Wang, Minu Pilvankar, Masoud Nickaeen, Steven Hansel, and Aleksander S Popel. Using quantitative systems pharmacology modeling to optimize combination therapy of anti-PD-L1 checkpoint inhibitor and T cell engager. *Frontiers in Pharmacology* **14** (2023), p. 1163432.
- [8] Yu Zhang, Aleksander S Popel, and Hojjat Bazzazi. Combining multikinase tyrosine kinase inhibitors targeting the vascular endothelial growth factor and cluster of differentiation 47 signaling pathways is predicted to increase the efficacy of antiangiogenic combination therapies. *ACS Pharmacology & Translational Science* **6.5** (2023), pp. 710–726.
- [9] Yu Zhang, Hanwen Wang, Rebeca Hannah M Oliveira, Chen Zhao, and Aleksander S Popel. Systems biology of angiogenesis signaling: Computational models and omics. *WIREs mechanisms of disease* **14.4** (2022), e1550.
- [10] Chen Zhao, Joshua L Heuslein, Yu Zhang, Brian H Annex, and Aleksander S Popel. Dynamic multiscale regulation of perfusion recovery in experimental peripheral arterial disease: a mechanistic computational model. *Basic to Translational Science* **7.1** (2022), pp. 28–50.
- [11] Yu Zhang, Christopher D Kontos, Brian H Annex, and Aleksander S Popel. A systems biology model of junctional localization and downstream signaling of the Ang–Tie signaling pathway. *npj Systems Biology and Applications* **7.1** (2021), p. 34.
- [12] Mohammad Jafarnejad, Richard J Sové, Ludmila Danilova, Adam C Mirando, Yu Zhang, Mark Yarchoan, Phuoc T Tran, Niranjana B Pandey, Elana J Fertig, and Aleksander S Popel. Mechanistically detailed systems biology modeling of the HGF/Met pathway in hepatocellular carcinoma. *NPJ systems biology and applications* **5.1** (2019), p. 29.
- [13] Yu Zhang, Christopher D Kontos, Brian H Annex, and Aleksander S Popel. Angiopoietin-Tie signaling pathway in endothelial cells: a computational model. *Iscience* **20** (2019), pp. 497–511.
- [14] Chen Zhao, Yu Zhang, and Aleksander S Popel. Mechanistic computational models of microRNA-mediated signaling networks in human diseases. *International Journal of Molecular Sciences* **20.2** (2019), p. 421.

- [15] Hojjat Bazzazi, Yu Zhang, Mohammad Jafarnejad, Jeffrey S Isenberg, Brian H Annex, and Aleksander S Popel. Computer simulation of TSP1 inhibition of VEGF–akt–eNOS: an angiogenesis triple threat. *Frontiers in physiology* **9** (2018), p. 644.
- [16] Hojjat Bazzazi, Yu Zhang, Mohammad Jafarnejad, and Aleksander S Popel. Computational modeling of synergistic interaction between $\alpha V\beta 3$ integrin and VEGFR2 in endothelial cells: Implications for the mechanism of action of angiogenesis-modulating integrin-binding peptides. *Journal of theoretical biology* **455** (2018), pp. 212–221.
- [17] Yu Zhang, Hojjat Bazzazi, Raquel Lima e Silva, Niranjana B Pandey, Jordan J Green, Peter A Campochiaro, and Aleksander S Popel. Three-dimensional transport model for intravitreal and suprachoroidal drug injection. *Investigative Ophthalmology & Visual Science* **59.12** (2018), pp. 5266–5276.

Book chapters

- [1] Yu Zhang, Chen Zhao, and Aleksander S. Popel. “Systems biology modeling of endothelial cell and macrophage signaling in angiogenesis in human diseases”. *The Vasculome*. Academic Press, **2022**, pp. 163–172.

Manuscripts under review

- [1] Yu Zhang, Andrew Hennes, Aarti Krishnan, Satotaka Omori, Alicia Li, Leif Sieben, Ronak Desai, Seyed Majed Modaresi, Maxwell Z. Wilson, Felix Wong, and James J. Collins. Multitask learning enables discovery of nontoxic antibiotics. *Under Review* (2025).
- [2] Yu Zhang, Hanwen Wang, Natsuki Furukawa, Akash Patil, Theinmozhi Arulraj, and Aleksander S. Popel. Quantitative systems pharmacology model predicts enhanced antitumor efficacy of anti-CTLA-4 when combined with anti-PD-1 in syngeneic mouse model of breast cancer. *Under Review* (2025).
- [3] Melis N. Anahtar, Jacqueline A. Valeri, Seyed Majed Modaresi, Aarti Krishnan, Nina M. Donghia, Samantha G. Palace, Erica J. Zheng, Aakanksha Gulati, Alicia Jorgenson, Abidemi Junaid, Parijat Bandyopadhyay, Andreas Lutten, Krishna Suresh, Paige Edwards, Felix Wong, Yu Zhang, Danilo Ritz, Margaux Gaborieau, Edmund Loh, Massimiliano Gaetani, Marie-Stephanie Aschtgen, Amir Ata Saei, Yonatan H. Grad, Donald E. Ingber, and James J. Collins. Deep learning-enabled discovery of antibiotics effective against *Neisseria gonorrhoeae*. *Under Review* (2025).

Selected Presentations

ACS Spring 2025 – Multitask Learning Enables Discovery of Nontoxic Antibiotics	2025
<i>Session speaker (Symposium on AI in Drug Discovery), San Diego, CA</i>	
NAVBO In-Focus Webinar – Endothelial Regulation of Microvascular Permeability	2024
<i>Invited webinar speaker, Virtual</i>	
Foundations of Systems Biology in Engineering (FOSBE)	2022
<i>Session speaker, Boston, MA</i>	

Leadership, Service & Recognition

Organizing Committee	2024 – Present
<i>Broad Institute Machine Learning for Drug Discovery Symposium</i>	
Co-design the scientific program and recruit speakers for an annual symposium that bridges AI and drug discovery, shaping the community’s research agenda.	
Review Editor	2023 – Present
<i>Frontiers in Pharmacology</i>	
Lead Teaching Assistant	2016, 2020
<i>JHU: Models and Simulations (EN.580.223)</i>	<i>Baltimore, MD</i>
Benjamin Zweifach Student Travel Award	2022
<i>The Microcirculatory Society (Travel Award)</i>	
Richard J. Johns Award for Outstanding Academic Achievement	2017
<i>JHU Dept. of Biomedical Engineering</i>	<i>Baltimore, MD</i>