Harrison Specht

Graduate Student, Bioengineering, Northeastern University 360 Huntington Ave, Boston, MA 02110

360 Huntington Ave, Boston, MA 02110

Specht.h@northeastern.edu Sharrisonspecht.com | Updated: February 23, 2021

EDUCATION Northeastern University, Ph.D. Bioengineering, Graduate Student expected 2021 Cornell University, B.A. Chemistry, magna cum laude 2014 RESEARCH EXPERIENCE *Northeastern University* Slavov Lab, Bioengineering & Barnett Institute, Graduate Student 2016-Merck, Sharpe & Dohme Corporation Spring 2020 Merck Exploratory Sciences Division, Coop Broad Institute of MIT and Harvard Proteomics Platform, Research Associate 2014-2016 Cornell University, Ithaca, NY Petersen Lab, Department of Chemistry, Research Assistant 2012-2014 Zinder Lab, Department of Microbiology, Research Assistant 2010-2011 The Triple Helix, magazine, Executive Editor-in-chief 2013 - 2014Bermuda Institute for Ocean Sciences National Science Foundation REU Fellow 2013 Vertex Pharmaceuticals, Boston, MA Drug Metabolism and Pharmacokinetics, Internship 2013 FELLOWSHIPS AND AWARDS 2020 Rising Stars in Proteomics and Metabolomics, Journal of Proteome Research. https://pubs.acs.org/doi/full/10.1021/acs.jproteome.oco1026 2019 2nd place in Chemical Biology, Royal Society of Chemistry Twitter Poster Conference 2016- College of Engineering Dean's Distinguished Fellowship, Northeastern University

2013 Best Oral Presentation, Bermuda Institute for Ocean Science NSF REU Program

2013 National Science Foundation REU Fellowship, Bermuda Institute for Ocean Sciences

1/4

PUBLICATIONS

Specht, H., Emmott, E., Petelski, A.A. et al. Single-cell proteomic and transcriptomic analysis of macrophage heterogeneity using SCoPE₂. Genome Biol 22, 50 (2021). https://doi.org/10.1186/s13059-021-02267-5

Specht, H., Slavov, N., 2021. Optimizing accuracy and depth of protein quantification in experiments using isobaric carriers. J. Proteome Res. 20, 880-887. https://doi.org/10.1021/acs.jproteome.ocoo675

Huffman, R.G., Chen, A., **Specht, H.**, Slavov, N., 2019. DO-MS: Data-Driven Optimization of Mass Spectrometry Methods. J. Proteome Res. 18, 2493–2500. https://doi.org/10.1021/acs.jproteome.gbo0039

Specht, H., Harmange, G., Perlman, D.H., Emmott, E., Niziolek, Z., Budnik, B., Slavov, N., 2018. Automated sample preparation for high-throughput single-cell proteomics. bioRxiv 399774. https://doi.org/10.1101/399774

Specht, H., Slavov, N., 2018. Transformative Opportunities for Single-Cell Proteomics. J. Proteome Res. 17, 2565–2571. https://doi.org/10.1021/acs.jproteome.8b00257

Khajuria, R.K., Munschauer, M., Ulirsch, J.C., Fiorini, C., Ludwig, L.S., McFarland, S.K., Abdulhay, N.J., **Specht, H.**, Keshishian, H., Mani, D.R., Jovanovic, M., Ellis, S.R., Fulco, C.P., Engreitz, J.M., Schütz, S., Lian, J., Gripp, K.W., Weinberg, O.K., Pinkus, G.S., Gehrke, L., Regev, A., Lander, E.S., Gazda, H.T., Lee, W.Y., Panse, V.G., Carr, S.A., Sankaran, V.G., 2018. Ribosome Levels Selectively Regulate Translation and Lineage Commitment in Human Hematopoiesis. Cell 173, 90-103.e19. https://doi.org/10.1016/j.cell.2018.02.036

Keshishian, H., Burgess, M.W., **Specht, H.**, Wallace, L., Clauser, K.R., Gillette, M.A., Carr, S.A., 2017. Quantitative, multiplexed workflow for deep analysis of human blood plasma and biomarker discovery by mass spectrometry. Nat Protoc 12, 1683–1701. https://doi.org/10.1038/nprot.2017.054

TALKS

2020. **Harrison Specht**, Edward Emmott, Aleksandra A. Petelski, R. Gray Huffman, David H. Perlman, Marco Serra, Peter Kharchenko, Antonius Koller, Nikolai Slavov. "How to perform quantitative single cell proteomics with SCoPE2." GenomeWeb Webinar Series. *Invited*

2020. **Harrison Specht**, Edward Emmott, Aleksandra A. Petelski, R. Gray Huffman, David H. Perlman, Marco Serra, Peter Kharchenko, Antonius Koller, Nikolai Slavov. "How to perform quantitative single cell proteomics with SCoPE2." Association of Biomolecular Resource Facilities (ABRF) 2020 Annual Meeting. *Invited*

2019. **Harrison Specht**, Edward Emmott, Aleksandra A. Petelski, R. Gray Huffman, David H. Perlman, Antonius Koller, Nikolai Slavov. "Design of single cell proteomics experiments." Single Cell Proteomics Conference. Boston, MA. 2019.

2019. **Harrison Specht**, Nikolai Slavov. Quantifying proteins by mass spectrometry. Models, Inference, and Algorithms Seminar. Broad Institutde of MIT and Harvard. *Chalk-talk*

2018. **Harrison Specht**, Guillaume Harmange, David H Perlman, Edward Emmott, Zachary Niziolek, Bogdan Budnik, Nikolai Slavov. "Automated sample preparation for high-throughput single-cell proteomics." HUPO 2018.

2 2/4

2018. **Harrison Specht**, Guillaume Harmange, David H Perlman, Edward Emmott, Zachary Niziolek, Bogdan Budnik, Nikolai Slavov. "Automated sample preparation for high-throughput single-cell proteomics." Single Cell Proteomics Conference. Boston, MA. 2018.

POSTERS

2019. **Harrison Specht**. "Automated sample preparation for high-throughput single-cell proteomics." Royal Society of Chemistry Twitter Poster Conference. *Twitter poster presenter* Awarded 2nd place in Chemical Biology category.

2018. Hasmik Keshishian, Luke Wallace, **Harrison Specht**, Judit Jan-Valbuena, Rob McDonald, Dale Petterson, Eric Kuhn, Michael Burgess, D. R. Mani, Tomas Rejtar, Javad Golji, Karen Wang, William Sellers, Steven A. Carr. "SigPath300: A high throughput MS-based assay to quantify over 300 phosphosites of known biological relevance in cells and tissues." American Society for Mass Spectrometry Annual Meeting. *Contributor*

2017. Bogdan Budnik, Ezra Levy, **Harrison Specht**, Nikolai Slavov. "Exploring cell division dynamics across single cell proteomes." Cold Spring Harbor Laboratories: Single Cell Analyses. *Poster Presenter*

2017. **Harrison Specht**, Ezra Levy, Bogdan Budnik, Nikolai Slavov. "Mass-spectrometry of single mammalian cells quantifies proteome heterogeneity during cell differentiation." American Society for Mass Spectrometry Annual Meeting. *Poster Presenter*

2015. **Harrison Specht**, Hasmik Keshishian, Rajiv K. Khajuria, et al. "Addressing the challenge of ribosomal protein stoichiometry by proteomics." Broad Institute of MIT and Harvard Retreat. *Poster Presenter*

2013. **Harrison Specht** and Andrew Peters "Investigating North Atlantic Tar in Bermuda: the hydrocarbon composition of North Atlantic pelagic tar and their hydrocarbon leaching into seawater." BIOS NEF REU Program. *Presenter*

PROFESSIONAL AFFILIATIONS, TRAINING & SERVICE

- Volunteer moderator for Single Cell Proteomics Conference (Zoom) 2020
- Volunteer for Single Cell Proteomics Conference 2018, 2019
- Northeastern Univeristy Graduate Student Liason, 2018-2019
- Member of Northeastern University Fencing Club
- Member of Barnett Institute for Chemical and Biological Analysis
- eLife Ambassador, 2018
- ASAPbio Ambassador, 2018
- American Society for Mass Spectrometry
- May Institute on computation and statistics for mass spectrometry and proteomics, 2016
- Lab preprint journal club founder

TECHNIQUES

Coding and Software: R, Shiny, MS-Fragger, DIA-NN, SpectroNaut, MaxQuant, Skyline

Experimental: Single-cell proteomics, triple-quad and orbitrap-based mass spectrometry proteomics, data dependent (DDA) and data independant (DIA) acquisition, protein and peptide N-terminomics, phosphoproteomics, antibody and chemical enrichment of proteins and peptides, reverse-phase and ERLIC liquid chromatography, robotic sample preparation, cell culture, flow cytometry

3

REFERENCES

Available upon request

4