DAVID N.P. KOPPSTEIN

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Education

Massachusetts Institute of Technology Ph.D, Biology

Cambridge, MA 2015

Yale University

New Haven, CT

B.S., Molecular Biophysics and Biochemistry

2008

Experience

Bioinformatics Core, MPI for Immunobiology and Epigenetics **Staff Bioinformatician**

Freiburg, Germany

August '22 – present

Developed pipelines for demultiplexing and processing high-throughput sequencing data for end users. Spearheaded projects on transcriptome isoform annotation and chromosomal conformation capture using long reads and virus detection in single-cells.

Nikolaus Rajewsky Lab, Max Delbrück Center for Molecular Medicine

Berlin, Germany

Postdoctoral Researcher

June '18 – June '22

Conceived, wrote, and executed a Chan-Zuckerberg Initiative Neurodegeneration Grant to use 10X Visium spatial and Nanopore long read sequencing in tandem to analyze aberrant RNA isoforms in neuromuscular organoids models of Amyotrophic Lateral Sclerosis. Conceived, wrote, and executed a grant from the Berlin Institute of Health to research the effects of COVID-19 patient serum the endothelial transcriptome. Wrote and executed a grant from the Bundesministerium für Bildung und Forschung to explore the role of miR-155 and vault RNAs in SARS-CoV-2 infection. Performed sample preparation, RNA-seq library construction, and downstream bioinformatic analysis for these and other projects. Trained one Masters student.

Viral Immunology Systems Program, Kirby Institute, UNSW

Sydney, Australia

Postdoctoral Researcher

Apr '17 – April '18

Developed VDJPuzzle, a computational method for analyzing single-cell transcriptomics data of immune cells. Used this data and other methods to analyze single-cell transcriptomes of rogue B cells in Sjögren's disease. Performed single-cell RNA-seq using 10X Chromium and downstream bioinformatic analysis. Mentored two Ph.D students, a Masters student, and an Honors student.

Juno Therapeutics

SEATTLE, WA, USA

Data Scientist

Jan '16 – Mar '17

Principal architect of the immunosequencing bioinformatics pipeline. Scaled pipeline to the AWS cloud using CfnCluster and Snakemake. Codiscovered effective TCR binders for HPV E6 and E7, resulting in a patent. Analyzed and communicated immunosequencing analyses for internal projects and external academic collaborators. Conducted analyses for internal RNA-seq and vector integration profiling projects. Served as interim manager of the data science team during the acquisition.

AbVitro

BOSTON, MA, USA

Data Scientist

Mar 15 – Dec 15

Principal architect of the immunosequencing pipeline for single-cell immunosequencing. Mined receptor sequences from tumor-infiltrating lymphocytes and developed automated data visualization strategies to nominate candidates for screening. Communicated with external academic and industry collaborators. (*Acquired by Juno Therapeutics*).

David Bartel Lab, MIT

BOSTON, MA, USA

Graduate Student

May 10 - Mar 15

Designed, implemented, and analyzed high-throughput sequencing experiments using the Illumina platform to investigate influenza's cap-snatching repertoire, discovering that small nuclear RNAs are the primary targets of this process. Utilized 3P-seq to precisely examine alternative cleavage and polyadenylation in diverse genetic backgrounds and model organisms in order to update annotations and improve microRNA targeting models.

Joan Steitz Lab, Yale

New Haven, CT, USA

Undergraduate Student

May '06 – Apr '08

Investigated the function of Herpesvirus saimiri Small U RNAs (HSURs) by cloning aptamers, pulldowns, and mass spectrometry.

Skills

Wet lab

Extensive experience with NGS library preparation and custom design: 3P-Seq, SMART-seq, high-throughput 5' RACE, and standard RNA-seq. Experience running single-cell 10X Chromium, 10X Visium spatial sequencing, and Nanopore long-read sequencing. Experience with molecular cloning, tissue culture, Northern/Western blots, protein purification, PCR/qPCR, microscopy, cryotome sectioning, and yeast genetics.

Dry lab

Fluent in Python and R. Expertise with modern bioinformatics tools including samtools, STAR, featureCounts, edgeR, Picard, GSEA, etc. Experience with analysing single-cell and spatial sequencing data using scater, Seurat, napari, etc. Experience writing scalable computational pipelines using Snakemake and scaling to the cloud using AWS. Experience with statistical methods and machine learning techniques. Experience managing a team of computational biologists using agile methods.

Fellowships and Awards

- Juno CEO Award for Outstanding Performance, 2016
- NSF Graduate Research Fellowship Program, Honorable Mention, 2011
- Yale STARS II Research Fellowship, 2007-8

Grants

Wolfgang Kübler, Holger Gerhardt, Philipp Mertins, Nikolaus Rajewsky, Martin Witzenrath, **David Koppstein**, Laura Michalick, Anna Szymborska-Mell. The COVID-19 puzzle - The microvascular barrier as a mi ssing piece between coagulation and organ failure. Berlin Institute for Health, Vascular Biomedicine Focus Area Grant.

Mina Gouti and Nikolaus Rajewsky. Spatial Analysis of aberrant RNA isoforms in ALS neuromuscular organoids. Neurodegeneration Challenge Network Collaborative Pair Pilot Project Award. (Written on behalf of Nikolaus Rajewsky)

Nikolaus Rajewsky, Marcel Müller, and Markus Landthaler. Molecular exploration of SARS-CoV-2 host-virus interactions with targeted knockouts and pulldowns. Bundesministerium für Bildung und Forschung, COVID-19 awards. (Written on behalf of Nikolaus Rajewsky).

Presentations and Posters

Koppstein D, Rizzetto S, Samir J, Singh M, Reed JH, Cai CH, Lloyd AR, Eltahla AA, Goodnow CC, Luciani F. VDJPuzzle: A computational method for BCR and TCR reconstruction from single-cell sequencing data. Australian Bioinformatics And Computational Biology Society. Oral presentation delivered in Adelaide, Australia, November 2017.

Koppstein D Rizzetto S, Samir J, Singh M, Reed JH, Cai CH, Lloyd AR, Eltahla AA, Goodnow CC, Luciani F. VDJPuzzle: A computational method for BCR and TCR reconstruction from single-cell sequencing data. Australian Cellular Panomics Consortium. Poster presentation delivered in Melbourne, Australia, November 2017.

Koppstein D, Rizzetto S, Samir J, Singh M, Reed JH, Cai CH, Lloyd AR, Eltahla AA, Goodnow CC, Luciani F. VDJPuzzle: A computational method for BCR and TCR reconstruction from single-cell sequencing data. Australian Society for Immunology. Oral presentation delivered in Bowral, Australia, November 2017.

Koppstein D, Ashour J, Bartel D. Quantitative assessment of influenza's cap-snatching repertoire by RNA sequencing. RNA Society. Poster presentation delivered in Quebec City, Canada, June 2014.

Publications and Preprints

Bujanic L, Shevchuk O, von Kügelgen N, Kalinina A, Ludwik K, **Koppstein D**, Zerna N, Sickmann A, Chekulaeva M. The key features of SARS-CoV-2 leader and NSP1 required for viral escape of NSP1-mediated repression. 2022. *RNA. doi:* 10.1261/rna.079086.121.

Wyler M, Mösbauer K, Franke F, Diag A, Gottula LT, Arsie R, Klironomos F, **Koppstein D**, ..., Rajewsky N, Drosten C, Landthaler M. Transcriptomic profiling of SARS-CoV-2 infected human cell lines identifies HSP90 as target for COVID-19 therapy. 2021. *iScience. doi:* 10.1016/j.isci.2021.102151.

Singh M, Jackson KJL, Wang JJ, Schofield P, Field MA, **Koppstein D**, ..., Luciani F, Gordon TP, Goodnow CC, Reed JH. Lymphoma driver mutations in the pathogenic evolution of an iconic human autoantibody. 2020. *Cell. doi:10.1016/j.cell.2020.01.029*.

Rizzetto S, **Koppstein D**, Samir J, Singh M, Reed JH, Cai CH, Lloyd AR, Eltahla AA, Goodnow CC, Luciani F. B-cell receptor reconstruction from single-cell RNA-seq with VDJPuzzle. 2018. *Bioinformatics*. doi:10.1093/bioinformatics/bty203.

Grüning B, Dale R, Sjödin A, Chapman BA, Rowe J, Tomkins-Tinch CH, Valieris R, Koster J, **The Bioconda Team**. Bioconda: sustainable and comprehensive software distribution for the life sciences. 2018. *Nature Methods* 15, 475-76.

Grigaityte K, Carter JA, Goldfless SJ, Jefferey EW, Hause RJ, Jiang Y, **Koppstein D**, Briggs AW, Church GM, Vigneault F, Atwal GS. Single-cell sequencing reveals $\alpha\beta$ chain pairing shapes the T cell repertoire. 2017. *bioRxiv*.

Briggs AW, Goldfless SJ, Timberlake S, Belmont BJ, Clouser CR, **Koppstein D**, Sok D, Vander Heiden JA, Tamminen MV, Kleinstein SH, Burton DR, Church GM, Vigneault F. 2017. Tumor-infiltrating immune repertoires captured by single-cell barcoding in emulsion. *bioRxiv*.

Koppstein D, Ashour J, and Bartel DP. Sequencing the cap-snatching repertoire of H1N1 influenza provides insight into the mechanism of viral transcription initiation. 2015. *Nucleic Acids Research* 43(10), 5052-5064.

Hezroni H, **Koppstein D**, Schwartz M, Tabin CJ, Bartel DP, and Ulitsky I. 2015. Principles of long noncoding RNA evolution derived from direct comparison of transcriptomes in 14 vertebrates. *Cell Reports* 11(7), 1110-1122.

Nam J-W, Rissland OS, **Koppstein D**, Abreu-Goodger C, Jan CH, Agarwal V, Yildirim, Rodriguez A, and Bartel DP. 2014. Global Analyses of the Effect of Different Cellular Contexts on MicroRNA Targeting. *Molecular Cell.* 53, 1031-1043.

Ulitsky I, Shkumatava A, Jan C, Subtelny AO, **Koppstein D**, Bell G, Sive H, and Bartel DP. 2012. Extensive alternative polyadenylation during zebrafish development. *Genome Research*. 22(10):2054-66.

Agarwal A*, **Koppstein D***, Rozowsky J, Sboner A, Habegger L, Hillier LW, Sasidharan R, Reinke V, Waterston RH, and Gerstein M. 2010. Comparison and calibration of transcriptome data from RNA-Seq and tiling arrays. *BMC Genomics*. 11:383.

Mukhopadhyay J, Das K, Ismail S, **Koppstein D**, Jang M, Hudson B, Sarafianos S, Tuske S, Patel J, Jansen R, Irschik H, Arnold E, and Ebright RH. 2008. Myxopyronin, Corallopyronin, and Ripostatin Inhibit Transcription by Binding to the RNA Polymerase Switch Region. *Cell.* 135:295-307.

Extracurricular

- Instructor for NGSSchool seminar, "Introduction to Snakemake", 2020
- Science outreach with "Lange Nacht für Wissenschaft" (Long Night of Science), Berlin, 2019-2021
- Urban Cycling Skills and Safety Clinic Organizer, 2013-4
- Whitehead Partner for High School Science Teacher Outreach, 2012
- MIT Cycling Team Officer, 2012-2014

Languages

- English native speaker
- German B2 level

 $^{^*}$ Authors contributed equally to this publication.