# **DEVISHI KESAR**

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#### **EDUCATION**

## German Cancer Research Center(DKFZ)

Heidelberg, Germany

University of Heidelberg

Doctor of Natural Science

October 2026

• Relevant Skills: Multi-modal data integration, analysis of ChIP-seq, DNA methylation, copy number variation (CNV)

## Georgia Institute of Technology

Atlanta, GA, USA

Master's in Bioinformatics

December 2020

• Relevant Skills: Variant calling analysis, differential expression analysis, database systems management

#### Indraprastha Institute of Technology Delhi

Delhi, India

Bachelor of Technology (Honors) in Computer Science and Engineering Minor in Computational Biology

August 2019

• Relevant Skills: Machine Learning, statistical analysis, graph modeling, big data analysis

# **SKILLS**

Programming and Computational Skills: Python, Bash, R, Java, Scala, Regex, SQL, MATLAB Tools and Platforms: Git, Hive, Cytoscape, RStudio, PyMol, Databricks, AWS, Spark, Docker, Google Cloud Services

#### **PUBLICATIONS**

- Chaturantabut, S., Oliver, S., Frederick, D.T., Kim, J., Robinson, F.P., Sinopoli, A., Song, T.Y., He, Y., Chang, Y.C., Rodriguez, D.J., Chang, L., Kesar, D., Ching, M., Dzvurumi, R., Atari, A., Tseng, Y.Y., Bardeesy, N., Sellers, W.R., Identification of potent biparatopic antibodies targeting FGFR2 fusion driven cholangiocarcinoma. (2025). The Journal of Clinical Investigation
- Chang, L., Jung, N.Y., Atari, A., Rodriguez, D.J., Kesar, D., Song, T.Y., Rees, M.G., Ronan, M., Li, R., Ruiz, P., Chaturantabut, S., Ito, T., Tienen, L.M., Tseng, Y.Y., Roth, J., & Sellers, W.R. Systematic profiling of conditional pathway activation identifies context-dependent synthetic lethalities. (2023). Nature Genetics
- •Li, R., Klingbeil, O., Monducci, D., Young, M. J., Rodriguez, D. J., Bayyat, Z., Dempster, J. M., Kesar, D., Yang, X., Zamanighomi, M., Vakoc, C. R., Ito, T., & Sellers, W. R. Comparative optimization of combinatorial CRISPR screens. (2022). Nature Communications
- Mulvaney, K. M., Blomquist, C., Acharya, N., Li, R., Ranaghan, M. J., O'Keefe, M., Rodriguez, D. J., Young, M. J., Kesar, D., Pal, D., Stokes, M., Nelson, A. J., Jain, S. S., Yang, A., Mullin-Bernstein, Z., Columbus, J., Bozal, F. K., Skepner, A., & Sellers, W. R. Molecular basis for substrate recruitment to the PRMT5 methylosome. (2021). Molecular Cell.
- Mason, M.J., Schinke, C., Eng, C.L.P. et al. Multiple Myeloma DREAM Challenge reveals epigenetic regulator PHF19 as marker of aggressive disease. Leukemia (2020) - member of DREAM consortium
- Yang, et al. Assessment of the Limits of Predictability of Protein and Phosphorylation Levels in Cancer. (2020), Cell Systems member of NCI-CPTAC-DREAM consortium

## MANUSCRIPTS IN PREPARATION

- Knoll, N., Masser, S., Kesar, D., Bordas, B., Kania, N., Zagar, S., Rodriguez, D.J., Jen, J., He, Y., Li, G., Mensah, C., Chen, Z., Feng, B., Chokshi, M., Raghavan, S., Sellers, W.R. and Mulvaney, K.M., CRISPR-Drug Combinatorial Screening Identifies Effective Combination Treatments for MTAP null cancer
- Shirole, N., Kesar, D., Lee, Y., Goodale, A., Syamala, S., Kukreja, S., Li, R., Qiu, X., Yu, W., Goldman, S., Cejas, P., Long, H., Adelman, K., Doench, J., Sellers, W.R. and Kaelin, W., Sustaining Cyclin D1 Expression Renders Kidney Cancer Cells HIF2-Independent

#### RESEARCH EXPERIENCE

## Pediatric glioma research lab, DKFZ

#### Heidelberg, Germany

Graduate student

October 2023 - Present

- Processed ChIP-seq data to identify transcription factor binding patterns and histone modifications in pediatric gliomas
- Integrated RNA-seq and ChIP-seq data to link super-enhancer regions with gene expression
- Conducted transcription factor motif analysis to explore regulatory elements driving tumor-specific pathways
- Performed methylation and copy number variation analyses to uncover genetic and epigenetic changes in tumor subtypes
- Leveraged deep variational autoencoders and MOFA analysis to integrate multi-modal data, uncovering new relationships
- Constructed a gene regulatory network, uncovering novel biomarkers associated with different glioma subtypes

### Sellers Lab, Broad Institute of MIT and Harvard

### Cambridge, MA, USA

Associate Computational Biologist

April 2021 – September 2023

- Processed and analyzed data for CRISPR paralog redundancy screens to find actionable biomarkers
- Executed antibody docking and structure prediction for antibodies using modeling softwares
- Performed ChIPseq data analysis to look at TF motif enrichment and single nucleotide resolution cut profile
- Detected alternate splicing events helping suggest PBM interaction requirement for PRMT5-dependent splicing events
- Analyzed data from mass spectrometry experiments conducted to identify novel E3 substrates

#### Georgia Institute of Technology

#### Atlanta, GA, USA

Graduate Research Assistant

August 2019- December 2020

- Benchmarked variant calling pipelines by comparison of performance of mappers and variant callers
- Cataloged genomic variants from Colombian populations and characterized their predicted functional impact
- Identified single variant disease associations using best practices from the American College of Medical Genetics
- Evaluated distribution of health-related genetic variants among populations from Colombia in light of their ancestry

#### **EMD Sereno**

# Billerica, MA, USA

Bioinformatics Summer Intern

*May* 2020 – *July* 2020

- Developed Rshiny app to share isolated immune cell RNAseq from bulk breast cancer tumors with biological experts
- Discovered novel myeloid drug targets by leveraging differential expression and gene signature analysis
- Drove bioinformatics discovery by frequently presenting results to key stakeholders across multiple departments
- Identified actionable biomarkers supporting therapeutic remediation of immuno-oncology and autoimmune disease

## POSTER PRESENTATIONS

- Kesar, D., Keck, M.K., Joshi, P., Okonechnikov, K., Autry, R.J., T.W. Jones, D.T.W., Integrated (epi)genomic atlas for identification of biomarkers in pediatric gliomas, KiTZ retreat 2024
- Kesar, D. Colombian Genome Diversity Project, GT@SC20 HPC Poster Show Proceedings
- Kesar, D. The discovery of context-specific paralog dependencies in cancer, Broad Retreat 2022

#### **AWARDS**

- Computational Biology Research Award Spring 2020 and Fall 2020 Georgia Tech
- Dean Academic Excellence Award IIIT Delhi 2018
- RECOMB/ISCB Conference Travel Grant 2017

# TEACHING EXPERIENCE

• Bulk RNA-seq computational analysis

• Introduction to quantitative biology

Instructor: Dr. Subhadip Raychaudhuri

Nov 2024-Present Jan 2018-Apr 2018

# LEADERSHIP AND VOLUNTEER EXPERIENCE

- Member, Social events team, DKFZ, Germany 2023-25
- Organizing team, Broad Retreat 2022, Broad Institute, USA 2021-22
- Volunteer, Charles river cleanup, The Charles River Conservancy, MA, USA, 2021-22
- Volunteer, GP Covid testing lab, Broad Institute, MA, USA, 2022

# **COURSES AND CERTIFICATIONS**

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Research Management Training for Early-Stage-Researchers	Apr'25
CRUK Children's Brain Tumor Centre of Excellence Summer School	Jul'24
• Cancer Genomics and Precision Oncology, online course by Harvard Medical School	Mar'22
• Immuno-oncology, online course by Harvard Medical School	Mar'22
Precision medicine, online course by University of Geneva	Feb'21
• Starting single cell analysis, online course by EMBL-EBI	Jun'20
• Introduction to breast cancer, online course by Yale University	Jun'19
Computer science and automation, Summer School   IISc Bangalore	Jul'18
Summer school in Bioinformatics   NCBS	Jun'17