

EDUCATION

- **Middle East Technical University** Ankara, TURKEY
PhD in Computer Engineering; CGPA: 3.21/4.00 – GPA: 3.75/4.00 2016 – Present
 - Network modeling and analysis of the temporal signaling networks for ovarian cancer patients when there is a delay in freezing tumor tissue after the removal (ongoing study).
- **Middle East Technical University** Ankara, TURKEY
MSc in Bioinformatics; CGPA: 3.86/4.00 (95.80/100) – GPA: 3.83/4.00 (94.90/100) 2016
 - Network modeling and analysis of the temporal signaling network in *Salmonella*-infected human cells (MSc thesis study).
- **Middle East Technical University** Ankara, TURKEY
BSc in Biology; CGPA: 3.32/4.00 (83.20/100) – GPA: 3.75/4.00 (92.50/100) 2014
 - Developing a web-based software that constructs functional genetic devices with the input and output parameters which are defined as the activator of promoters and the production of coding sequences, respectively.
 - Design of a biosensor that can detect MRSA (Methicillin-resistant *S. aureus*).
- **Kirami Refia Alemdaroglu High School** Ankara, TURKEY
Mathematics and Science; Diploma Grade: 83/100 2009

EXPERIENCE

- **Seven Bridges Genomics, Inc.** Ankara, TURKEY
Bioinformatics Analyst Dec. 2016 – Present
 - Dockerizing and optimizing various bioinformatics tools for cloud computing on Seven Bridges Platform and Cancer Genomics Cloud.
 - Researching and developing bioinformatics algorithms and tools for next-generation sequencing (NGS) data analysis.
- **School of Informatics and Computing, IUPUI** Indianapolis, IN, USA
Bioinformatics Research Scholar Oct. 2015 – Jun. 2016
 - Developing a Python package and a React application for systematic identification and comparison of processes, phenotypes and diseases associated with RNA-binding proteins from condition-specific CLIP-seq profiles.
 - Transcriptome analysis of developing mouse lens tissue including novel transcript discovery and alternative splicing alterations.
 - Developing a database of transcriptome profiles encompassing known and novel transcripts across multiple developmental stages in eye tissues in mouse; the data have been curated from Sequence Read Archive (SRA) and European Nucleotide Archive (ENA) and quality check, alignment, transcript discovery and transcript quantification have been performed.
 - Developing a web resource for prognostic analyses of exons across human cancers using clinical transcriptomes obtained from The Cancer Genome Atlas (TCGA).
- **Evias Pharmaceutical R&D, Co., Ltd.** Ankara, TURKEY
Software Specialist Feb. 2015 – Oct. 2015
 - Developing a cloud-based (AWS EC2) high performance virtual screening for drug discovery web service by integrating different tools such as JSmol, JSME, Google Charts, Autodock 4, Autodock Vina, MolPort Web Services using Django (Python) web framework.
- **Atlas Biotechnologies, Co., Ltd.** Ankara, TURKEY
Intern May 2014 – Feb. 2015

- Foreign relations with biotechnology product suppliers, giving orders and tracking the process including customs regulations.
- Detecting possible mutations in sequences obtained from ABI format files by blasting against reference genomes database or aligning to the corresponding reference genome.

• **Bioinformatics Department, Maastricht University** Maastricht, THE NETHERLANDS
Intern Jun. 2013 – Sep. 2013

- Analysis of the dataset on four breast cancer cell lines to build network models that represent the active cell signaling pathways in breast cancer.

• **Bioinformatics Unit, German Cancer Research Center (DKFZ)** Heidelberg, GERMANY
Intern Jun. 2012 – Sep. 2012

- Contaminant detection and identification in Next-Generation Sequencing data by developing a pipeline which extracts the unmapped reads and compares the possible contaminant genomes in different databases using Perl programming language.

PUBLICATIONS

- **Budak, G.**, Dash, S., Srivastava, R., Lachke, S. A., & Janga, S. C. (2018). Express: A database of transcriptome profiles encompassing known and novel transcripts across multiple development stages in eye tissues. *Experimental eye research*.
<https://www.ncbi.nlm.nih.gov/pubmed/29337142>
- Srivastava, R., **Budak, G.**, Dash, S., Lachke, S. A., & Janga, S. C. (2017). Transcriptome analysis of developing lens reveals abundance of novel transcripts and extensive splicing alterations. *Scientific Reports*, 7(1), 11572.
<https://www.ncbi.nlm.nih.gov/pubmed/28912564>
- **Budak, G.**, Srivastava, R., & Janga, S. C. (2017). Seten: A tool for systematic identification and comparison of processes, phenotypes and diseases associated with RNA-binding proteins from condition-specific CLIP-seq profiles. *RNA*, rna-059089.
<https://www.ncbi.nlm.nih.gov/pubmed/28336542>
- Hashemikhabir, S., **Budak, G.**, & Janga, S. C. (2016). ExSurv: A Web Resource for Prognostic Analyses of Exons Across Human Cancers Using Clinical Transcriptomes. *Cancer Informatics*, 15(Suppl 2), 17.
<https://www.ncbi.nlm.nih.gov/pubmed/27528797>
- Hill, S. M., Heiser, L. M., Cokelaer, T., Unger, M., Nesser, N. K., Carlin, D. E., **The HPN-DREAM Consortium**, ... & Graim, K. (2016). Inferring causal molecular networks: empirical assessment through a community-based effort. *Nature methods*, 13(4), 310-318.
<https://www.ncbi.nlm.nih.gov/pubmed/26901648>
- **G. Budak**, O. Eren-Ozsoy, Y. Aydin-Son, T. Can, and N. Tuncbag, Reconstruction of the temporal signaling network in *Salmonella*-infected human cells, *Front. Microbiol.*, 6:00730, 2015.
doi:10.3389/fmicb.2015.00730.
<https://www.ncbi.nlm.nih.gov/pubmed/26257716>

ORAL PRESENTATIONS

- 2017 – Oral presentation titled "Identification of the Ischemic Pathway Level Changes by Integrating Temporal Phosphoproteome in Ovarian Cancer" at 5th International Congress of the Molecular Biology Association of Turkey / Bogazici University, Istanbul, TURKEY.
- 2012 – Oral presentation titled "METU Biology and Genetics Student Club Introduction" at FBI - DIYbio Outreach Workshop / Marriott Hotel, Walnut Creek, CA, USA.

POSTER PRESENTATIONS

- 2016 – Poster presentation titled "Seten: A tool for systematic identification and comparison of processes, phenotypes and diseases associated with RNA-binding proteins from condition-specific CLIP-seq profiles" at GLBIO/CCBC Great Lakes Bioinformatics and the Canadian Computational Biology Conference 2016 / University of Toronto, Toronto, CANADA.
Published on F1000Research: <https://f1000research.com/posters/5-2586>
- 2015 – Poster presentation titled "Reconstruction of the temporal signaling network in Salmonella-infected human cells" at ISMB/ECCB 2015, Intelligent Systems for Molecular Biology / European Conference on Computational Biology 2015 / The Convention Centre Dublin, Dublin, IRELAND.
Published on F1000Research: <http://f1000research.com/posters/4-292>
- 2013 – Poster presentation titled "CONTAMINATION: detect contaminating sequences in high-throughput sequencing data" at ISMB/ECCB 2013, Intelligent Systems for Molecular Biology / European Conference on Computational Biology 2013 / Messe Berlin, Berlin, GERMANY (contributed study was presented by co-authors).
Published on F1000Research: <http://f1000research.com/posters/1094020>
- 2012 – Paper presentation titled "M4B: A Novel Method for Designing and Ordering Genetic Constructs" at HIBIT 2012, 7th International Symposium on Health Informatics and Bioinformatics / Cappadocia, Nevsehir, TURKEY (contributed study was presented by co-authors).
Published on IEEE Xplore: <http://ieeexplore.ieee.org/xpl/articleDetails.jsp?arnumber=6209054>

SKILLS

- **Programming:** Python, R, JavaScript, C/C++, Java, PHP, Perl, HTML, CSS, MySQL, SQLite, NoSQL (MongoDB), L^AT_EX.
- **Bioinformatics tools:** NGS data QC (FastQC, Trim Galore!, Cutadapt), data manipulation (SAMTools), data analysis (BLAST, BWA, HISAT, GATK, MACS2, StringTie, rMATS, IGV, NetworkX, Cytoscape, Autodock4, PyMol, UCSC Chimera, Ensembl API, Rabix).
- **Technologies & Libraries:** Git, svn, AWS, Google Cloud, Wordpress, Django, Laravel, React, D3.js, Twitter Bootstrap, BioPython, Numpy, Pandas, Scipy, Scikit-learn, Matplotlib, igraph, ggplot2, ComplexHeatmap.
- **Softwares/OSes:** RStudio, Postman, Android Studio, Adobe Photoshop/Illustrator, MS Office tools, MS Windows, Unix/GNU/Linux, macOS.

LANGUAGES

- Turkish (native)
- English (fluent) – METU EPE score: 90.0/100 (2017)
- French (beginner)

AWARDS AND CERTIFICATES

- 2017 – Best Poster Award at HIBIT 2017, 10th International Symposium on Health Informatics and Bioinformatics, by ISCB Student Council Regional Study Group Turkey.
- 2016 – Thesis of the Year by METU Prof. Dr. Mustafa N. Parlar Education and Research Foundation.
- 2015 – Travel funding to ISMB/ECCB 2015, Intelligent Systems for Molecular Biology / European Conference on Computational Biology 2015 by BMC Bioinformatics.
- Certificate of high honor / certificate of honor: 2012 – 2013 spring, 2013 – 2014 fall / 2010 – 2011 fall, 2011 – 2012 fall, 2011 – 2012 spring, 2012 – 2013 fall.
- 2011 – iGEM (International Genetically Engineered Machines) Competition World Championship "Software Division: Best Use of Registry" Award / Massachusetts Institute of Technology, Cambridge, MA, USA.
- 2011 – iGEM (International Genetically Engineered Machines) Competition European Jamboree Silver Medal / Vrije University, Amsterdam, THE NETHERLANDS.
- 2011 – Hacettepe University Technopolis Technovation Competition Mention Award / Hacettepe University, Ankara, TURKEY.

VOLUNTEERING

- 2015 – Lilly AMECA 2015, Lilly Africa, Middle East and Central Asia (AMECA) 2015 / Indianapolis, IN, USA.
- 2015 – ISMB/ECCB 2015, Intelligent Systems for Molecular Biology / European Conference on Computational Biology 2015 / Dublin, IRELAND.
- 2013 – TEDxMaastricht / Maastricht, THE NETHERLANDS.
- 2010 – 2014: Organizing various scientific conferences on biodiversity, evolutionary biology and many other fields in biology as a member of METU Biology and Genetics Student Club / Ankara, TURKEY.

MEMBERSHIPS

- International Society for Computational Biology, ISCB (as a member since 2015).
- Tree of Evolution Movement (as the co-founder and developer since November, 2010)
<http://www.evrimagaci.org>.
- METU Biology and Genetics Student Club (as a member since September, 2009, as a member of administrative board between September, 2010 – February, 2011, as the head of the administrative board between February, 2011 – June, 2012, as a member of supervising council between June, 2012 – January, 2014).

INTERESTS AND HOBBIES

- Biology, molecular biology, genetics, computer science, statistics, bioinformatics, coding, programming, website development.
- Blogging on bioinformatics and English language; playing guitar; doing sports; playing basketball and football, volunteering.

REFERENCES

- Assoc. Prof. Dr. Nurcan TUNCBAG
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