

Ege Ulgen MD, PhD

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Genomics England Limited



he/him



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About me ——

MD-PhD professional in bioinformatics with a medical degree from Acibadem University in 2015, postdoctoral experience at Yale University in 2016, followed by a PhD in Biostatistics and Bioinformatics in 2022. Currently a Bioinformatics Engineer at Genomics England, where I focus on creating and enhancing software for the cancer genomic medicine service.

Education

2009-2015 M.D., summa cum laude

2016-2022 Ph.D., Biostatistics and Bioinformatics

Acibadem University

Acibadem University

[Experience]

tribble(Experience, ~ Year, ~ Institution, ~Description, "Bioinformatics Engineer", "2023-present", "Genomics England", "Developing features for and maintaining software for the cancer genomic medicine service.", "Bioinformatics Specialist", "2018-2022", "Epigenetiks", "Worked on developing and maintaining multiple -omics analysis pipelines for the local medical diagnostics company. Responsibilities included: implementation, continuous development and maintenance of whole exome sequencing analysis pipeline for aiding the diagnosis of undiagnosed subjects with rare diseases, and whole exome sequencing analysis pipeline for cancer diagnostics and prognostics.", "Visiting Researcher", "2018", "Akalin Lab, Max-Delbrück-Centrum für Molekulare Medizin", "Worked on a project on the effects of "doublet cells" on single-cell RNAseq analysis. Using a real data set and simulations, I demonstrated that doublets cause pseudo-paths (false patterns of differentiation), negatively affect discrimination between different cell types and have a negative effect on imputation results.", "Bioinformatics Consultant", "2015-2022", "NeuroOncology Technologies", "Developed and maintained the Whole Exome Sequencing pipeline for the local medical diagnostics company based in Istanbul-Turkey. The pipeline allows for the identification of clinically-relevant genomics findings in brain tumors, especially regarding gliomas", "Research Internships", "2011-2014", "Yale University School of Medicine", "Research in neurogenetics, particularly malformations of cortical development, utilizing molecular techniques to investigate the underlying genetic/molecular mechanisms, analyzing Whole Exome Sequencing results and verifying relevant findings. Research in brain tumours, performing functional in-vitro analyses to investigate the effect of a SNP in the pathogenesis of gliomas.", "Research Group Member", "2012-present", "ACU Brain Tumor Research Group", "Participate in research on brain tumors, particularly on gliomas. I assist in biostatistical as well as bioinformatics analyses, focusing on the integration of molecular biological and clinical data to improve diagnosis, prognosis and treatment of glioma.",) %>% detailed_entries(Experience, Year, Institution, Description)

Skills

Programming

python - 5/5 R - 5/5bash - 4/5 C++ - 2/5

Development

git - 4/5 Docker - 3/5 GitLab CI - 4/5 GitHub Actions - 4/5 PyCharm - 5/5

[Strengths]

Curiosity Proactively seeks innovative solutions.

Time Management Effectively prioritizes tasks, ensuring timely completion of milestones.

Quick Learner Adapts to new challenges swiftly.

Software

| 2.3.0 | pathfindR. Enrichment Analysis Utilizing Active Subnetworks |
|-------|-------------------------------------------------------------------------|
| 1.0.1 | PANACEA. Personalized Network-Based Anti-Cancer Therapy |
| 0.4.1 | Evaluation driveR. Prioritizing Cancer Driver Genes Using Genomics Data |

2.0.0 pathfindR.data. Data Package for 'pathfindR'

Certification

| 2016 | Certificate in Machine Learning. Grade: 94.8% – cours- |
|------|--------------------------------------------------------------------------------------------------------------------------------------|
| 2018 | era.org/verify/7QETTCDWHXHP Certificate in Finding Hidden Messages in DNA (Bioinformatics I) (with Honors). Grade: 96.0% – cours- |
| | era.org/verify/B5Y2QZ4AWTCL |
| 2019 | Certificate in Genome Sequencing (Bioinformatics II) (with Hon- |
| | ors). Grade: 99.0% – coursera.org/verify/242DMGMNGFXA |
| 2019 | Certificate in Comparing Genes, Proteins, and Genomes |
| | (Bioinformatics III) (with Honors). Grade: 100% - cours- |
| | era.org/verify/N3B77QC9N7V6 |
| 2019 | Certificate in Molecular Evolution (Bioinformatics IV) (with Hon- |
| | ors). Grade: 100% – coursera.org/verify/TM3MWQPRPS47 |
| 2019 | Certificate in Genomic Data Science and Clustering (Bioin- |
| | formatics V) (with Honors). Grade: 94.0% - cours- |
| | era.org/verify/AGX8F2V8PU3G |
| 2019 | Certificate in Bioinformatics Capstone: Big Data in Biology. Grade: |
| | 100% – coursera.org/verify/9AKDLPVN9M2U |
| 2019 | Certificate in Bioinformatics Specialization. cours- |
| | era.org/verify/35KWGTCVTJX8 |

Publications

- 1. Kaya, D., Ülgen, E., Kocagöz, A., & Sezerman, O. (2023). A comparison of various feature extraction and machine learning methods for antimicrobial resistance prediction in streptococcus pneumoniae. *Frontiers in Antibiotics*.
- 2. Ulgen, E., Ozisik, O., & Sezerman, O. (2023). PANACEA: Network-based methods for pharmacotherapy prioritization in personalized oncology. *Bioinformatics*.
- 3. Levi, C., Uçal, Y., Planchon, S., Ülgen, E., Kumru, P., Ulutaş, P., Sezerman, U., & (2023). Proteome analysis of human and goat colostrum: A closer look at whey fractions. *Acibadem Saglik Bilimleri Dergisi*.
- 4. Cansu, A., Yasemin, U., Planchon, S., Ülgen, E., KUMRU, P., ULUTAŞ, P., & (2023). Proteome analysis of human and goat colostrum: A closer look at whey fractions. *Acıbadem Üniversitesi Sağlık Bilimleri Dergisi*.
- 5. Dogan, B., Gumusoglu, E., Ulgen, E., Sezerman, O., & Gunel, T. (2022). Integrated bioinformatics analysis of validated and circulating miRNAs in ovarian cancer. *Genomics & Informatics*.
- 6. Keleş, I., Günel, T., Özgör, B., Ülgen, E., Gümüşoğlu, E., Hosseini, M., & (2022). Gene pathway analysis of the endometrium at the start of the window of implantation in women with unexplained infertility and unexplained recurrent pregnancy loss: Is Human Fertility.
- 7. Yousef, M., Ülgen, E., & Sezerman, O. (2021). CogNet: Classification of gene expression data based on ranked active-subnetwork-oriented KEGG pathway enrichment analysis. *PeerJ Computer Science*.
- 8. Ülgen, E., & Sezerman, O. (2021). driveR: A novel method for prioritizing cancer driver genes using somatic genomics data. *BMC Bioinformatics*.
- 9. Albayrak, İ., Azhari, F., Çolak, E., Balcı, B., Ülgen, E., Sezerman, U., Baştu, E., & (2021). Endometrial gene expression profiling of recurrent implantation failure after in vitro fertilization. *Molecular Biology Reports*.
- Ülgen, E., Aras, F., Coşgun, E., Erşen-Danyeli, A., Dinçer, A., Usseli, M.,
 & (2021). Correlation of anatomical involvement patterns of insular gliomas with subnetworks of the limbic system. *Journal of Neurosurgery*.
- 11. Ülgen, E., Can, Ö., Bilguvar, K., Boylu, C. A., Yüksel, Ş. K., & (2021). Sequential filtering for clinically relevant variants as a method for clinical interpretation of whole exome sequencing findings in glioma. *BMC Medical Genomics*.
- 12. Everest, E., Ülgen, E., Uygunoglu, U., Tutuncu, M., Saip, S., Sezerman, O., & (2021). Investigation of multiple sclerosis-related pathways through the integration of genomic and proteomic data. *PeerJ*.
- 13. Keleş, I., Ülgen, E., Erkan, M., Çelik, S., Aydın, Y., Önem, A., Kandemir, H., & (2020). Comparison of endometrial prostanoid profiles in three infertile subgroups: The missing part of receptivity? *Fertility and Sterility*.

- 14. Ülgen, E., Karacan, S., Gerlevik, U., Can, Ö., Bilguvar, K., Oktay, Y., Akyerli, C. B., & (2020). Mutations and copy number alterations in IDH wild-type glioblastomas are shaped by different oncogenic mechanisms. *Biomedicines*.
- 15. Ozduman, K., Ulgen, E., Karacan, S., Gerlevik, U., Can, O., Bilguvar, K., & (2020). Mutations and copy number alterations in diffuse gliomas are shaped by different mechanisms. *European Journal of Cancer*.
- 16. Ulgen, E., Ozisik, O., & Sezerman, O. (2019). pathfindR: An r package for comprehensive identification of enriched pathways in omics data through active subnetworks. *Frontiers in Genetics*.
- 17. Bastu, E., Demiral, I., Gunel, T., Ulgen, E., Gumusoglu, E., Hosseini, M., & (2019). Potential marker pathways in the endometrium that may cause recurrent implantation failure. *Reproductive Sciences*.
- Ülgen, E., Bektaşoğlu, P., Sav, M., Can, Ö., Danyeli, A., Hızal, D., Pamir, M., & (2019). Meningiomas display a specific immunoexpression pattern in a rostrocaudal gradient: An analysis of 366 patients. World Neurosurgery.
- Ülgen, E., Can, Ö., Bilguvar, K., Oktay, Y., Akyerli, C., Danyeli, A., Yakıcıer, M., & (2019). Whole exome sequencing–based analysis to identify DNA damage repair deficiency as a major contributor to gliomagenesis in adult diffuse gliomas. *Journal of Neurosurgery*.
- 20. Sezerman, O., Ulgen, E., Seymen, N., & Durasi, I. (2019). Bioinformatics workflows for genomic variant discovery, interpretation and prioritization. *Bioinformatics Tools for Detection and Clinical Interpretation of Genomic*....
- 21. Ulgen, E., & Ozisik, O. (2019). pathfindR: An r package for comprehensive identification of enriched pathways in omics data through active subnetworks. Front genet 10: 858 59 dogan b, gumusoglu e, ulgen e *Genomics Inform*.
- 22. Akyerli, C., Yüksel, Ş., Can, Ö., Erson-Omay, E., Oktay, Y., Coşgun, E., & . . . (2018). Use of telomerase promoter mutations to mark specific molecular subsets with reciprocal clinical behavior in IDH mutant and IDH wild-type diffuse gliomas. *Journal of Neurosurgery*.
- 23. Ulgen, E., Ozisik, O., & Sezerman, O. (2018). pathfindR: An r package for pathway enrichment analysis utilizing active subnetworks. *BioRxiv*.
- 24. Bastu, E., Demiral, I., Ulgen, E., Erkan, M., Celik, S., Aydin, Y., Onem, A., & (2018). Lipidomic analysis reveals increased TXA2 presence in non-receptive endometrium of recurrent miscarriage and repeated implantation failure patients. *HUMAN REPRODUCTION*.
- 25. Dogan, M., Demiral, I., Akgun, E., Ulgen, E., Gurel, B., Sahin, B., Aytan, A., &.... (2018). Endometrial proteomic signature in recurrent implantation failure. *HUMAN REPRODUCTION*.
- 26. Demiral, I., Gumusoglu, E., Ulgen, E., Hosseini, M., Ozgor, B., Dogan, M., & (2018). Endometrial transcriptomic pathways analysis in recurrent miscarriages and unexplained infertility. *HUMAN REPRODUCTION*.
- 27. Ozduman, K., Ulgen, E., Can, O., Akyerli, C., & Pamir, M. (2018). Analysis of mutational processes in 23 adult hemispheric diffuse gliomas identifies DNA-damage repair deficiency as a major contributor to gliomagenesis. *JOURNAL OF NEUROSURGERY*.
- 28. Siva, A., Everest, E., Ülgen, E., UygunoÄŸlu, U., Tütüncü, M., Saip, S., & (2018). Identification of multiple sclerosis related pathways through genome-proteome correlations. *ACTRIMS Forum*.
- 29. Çağlayan, A., Sezer, R., Kaymakçalan, H., Ulgen, E., Yavuz, T., & (2017). ALPK3 gene mutation in a patient with congenital cardiomyopathy and dysmorphic features. *Molecular Case Studies*.
- 30. Ülgen, E., & Asar, Ö. (2017). Tekrarlı gözlem ve sağkalım verilerinin bileşik modellenmesi. *Türkiye Klinikleri Biyoistatistik*.
- 31. Demiral, I., Bastu, E., Gunel, T., Sezerman, U., Gumusoglu, E., Ulgen, E., & (2017). Endometrial gene expression in patients with recurrent implantation failure. *Fertility and Sterility*.
- 32. Bastu, E., Gunel, T., Sezerman, O., Demiral, I., Gumusoglu, E., Ulgen, E., & (2017). RAC1 signaling pathway is crucial for etiology of repeated implantation failure (RIF). *Fertility and Sterility*.

- 33. Oktay, Y., Ülgen, E., Can, Ö., Akyerli, C., Yüksel, Ş., Erdemgil, Y., Durası, İ., & (2016). IDH-mutant glioma specific association of rs55705857 located at 8q24. 21 involves MYC deregulation. *Scientific Reports*.
- 34. Per, H., Canpolat, M., Bayram, A., Ulgen, E., Baran, B., Kardas, F., Gumus, H., & (2015). Clinical, electrodiagnostic, and genetic features of tangier disease in an adolescent girl with presentation of peripheral neuropathy. *Neuropediatrics*.
- 35. Bayram, A., Per, H., Quon, J., Canpolat, M., Ülgen, E., Doğan, H., Gumus, H., & (2015). A rare case of congenital fibrosis of extraocular muscle type 1A due to KIF21A mutation with marcus gunn jaw-winking phenomenon. *European Journal of Paediatric Neurology*.